NUCLEOTIDE SEQUENCES FOR TRANSCRIPTIONAL REGULATION IN CORYNEBACTERIUM GLUTAMICUM

CROSS-REFERENCE TO RELATED APPLICATIONS

This application claims priority to U.S. Provisional Application 60/248,219, filed November 15, 2000, which is incorporated herein by reference in its entirety.

STATEMENT REGARDING FEDERALLY-SPONSORED $\label{eq:regarding} \textbf{RESEARCH AND DEVELOPMENT}$

Not applicable.

BACKGROUND OF THE INVENTION

Field of the Invention

[0001] The invention relates to the areas of microbial genetics and recombinant DNA technology. The invention provides DNA sequences, vectors, microorganisms, and methods useful for inducing and regulating the expression of genes, including those that are involved in amino acid biosynthesis, in bacterial cells.

Background Art

[0002] Coryneform bacteria are Gram-positive bacteria frequently used for industrial-scale production of amino acids, purines, and proteins. Although coryneform bacteria, particular Corynebacterium species, have been widely used for industrial purposes for many years, the techniques of molecular biology have only recently been employed to augment the usefulness of these organisms in the production of amino acids and other products.

[0003] One way to improve the productivity of a microbial strain is to increase the expression of genes that control the production of a metabolite. Increasing

expression of a gene can increase the activity of an enzyme that is encoded by that gene. Increasing enzyme activity can increase the rate of synthesis of the metabolic products made by the pathway to which that enzyme belongs. In some instances, increasing the rate of production of a metabolite can unbalance other cellular processes and inhibit growth of a microbial culture. The modified culture will make more product per cell, but will not be able to generate enough cells per volume to show an improvement over the parent strain in a fermentor.

[0004] Transcription is the process by which an RNA molecule is synthesized from a DNA template and occurs by the interaction of a multisubunit enzyme complex, known as RNA polymerase, with a DNA molecule. The RNA that is synthesized by this process ultimately directs the production of protein products within the cell. In general, the rate at which RNA is synthesized from DNA, i.e., the transcription rate, directly influences the level of synthesis of the

corresponding protein product.

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Promoters are DNA sequence elements that regulate the rate at which genes are transcribed. Promoters can influence transcription in a variety of ways. For example, some promoters direct the transcription of their associated genes at a constant rate regardless of the internal external cellular conditions. Such promoters are known as constitutive promoters. In many cases, however, a promoter will direct transcription of its associated gene only under very specific cellular conditions. For example, promoters that turn off gene expression during the growth phase of a microbial culture, but turn on gene expression after optimal growth has been achieved can be used to regulate genes that control production of a metabolite. The new strain will have the same growth pattern as the parent but produce more product per cell. This kind of modification can also improve titer (g product/liter) and yield (g product/g glucose). Nucleotide sequences have been identified that can be used to increase or decrease gene expression in Corynebacterium species. These regulatable promoters can increase or decrease the rate at which a gene is transcribed depending on the internal and/or the external cellular conditions. Frequently, the presence of a factor, known as an

inducer, can stimulate the rate of transcription from a promoter. Inducers can interact directly with molecules that, themselves, physically interact with the promoter or with DNA sequences in the vicinity of the promoter. Alternatively, the action of an inducer in stimulating transcription from a promoter may be indirect. Whereas inducers function to amplify the level of transcription from a promoter, there is a class of factors, known as suppressors, that reduce or inhibit transcription from a promoter. Like inducers, suppressors can exert their effects either directly or indirectly.

[0006] Besides regulation through inducers and suppressors, certain promoters are regulated by temperature. For instance, a the level of transcription from a promoter may be increased when cells harboring that promoter are grown at a temperature that is greater than the optimum or normal growth temperature for that cell type. Similarly, there are promoters that will enhance gene expression in cells grown at temperatures below the normal growth temperature.

[0007] Promoters are found naturally in wild-type cells where they regulate the expression of specific genes. Promoters, however, are also useful as tools of molecular biology in that they can be isolated from their normal cellular contexts and engineered to regulate the expression of virtually any gene.

[0008] The use of regulatory sequences from Escherichia coli to control the expression of reporter genes in Corynebacterium have been documented. The lacI^q repressor and the tac promoter/reporter genes from E. coli were on plasmids that replicate in Corynebacterium. See, e.g., Morinaga, Y. et al., J. Biotechnol. 5:305-312 (1987). In addition, Ben-Samoun et al., FEMS Microbiology Letters 174:125 (1999), which is incorporated herein by reference in its entirety, disclose the use of the E. coli araBAD promoter and the araC activator on a plasmid which replicates in Corynebacterium glutamicum cells to stimulate the expression of the GFPuv reporter gene only when L-arabinose is present in the growth

medium. The authors acknowledge, however, that the level of expression from the *araBAD* promoter in *C. glutamicum* is 6.5 fold lower than that which was observed in *E. coli*, the native species for the promoter.

[0009] U.S. Patent Nos. 5,693,781 and 5,762,299, each of which are incorporated herein by reference in their entireties, disclose the isolation of promoter sequences from the coryneform bacteria, Brevibacterium flavum. Sequences described in these patents were isolated on the basis of their ability to direct expression of a reporter gene in B. flavum at a level that was greater than the expression level observed in B. flavum with the synthetic tac promoter. Also disclosed in U.S. Patent Nos. 5,693,781 and 5,762,299, each of which are incorporated herein by reference in their entireties, are B. flavum promoters capable of expressing a reporter gene in B. flavum cells when grown in medium containing: (a) ethanol but not glucose, and vice versa; (b) glucose but not fructose; and (c) glucose but not casein hydrolysates/yeast extract/glucose, and vice versa. The novel promoter sequences disclosed in the present application are different from those described in U.S. Patent Nos. 5,693,781 and 5,762,299.

[0010] A limited number of C. glutamicum promoters have been described to date. For example, the C. glutamicum aceA promoter is disclosed in Wendisch et al., Arch Microbiol. 168:262 (1997) and in U.S. Patent No. 5,700,661 (where it is termed the isocitrate lyase promoter), each of which are incorporated herein by reference in their entireties. In both of these references, the aceA promoter was linked to a reporter gene in transformed C. glutamicum cells, and produced an extracellular protein, not a product of metabolic engineering. Expression of the reporter gene was found to be greater in C. glutamicum transformants that were grown in the presence of acetate than it was for transformants grown in the presence of glucose (Wendisch et al., Arch Microbiol. 168:262 (1997)) or sucrose (U.S. Patent No. 5,700,661).

[0011] Similarly, the aceB promoter from C. glutamicum is disclosed in Wendisch et al., Arch Microbiol. 168:262 (1997) and in U.S. Patent No. 5,965,391, which is incorporated herein by reference in its entirety. Both of these

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references describe transcriptional fusions consisting of the *aceB* promoter region linked to a reporter gene in *C. glutamicum* transformed cells. Expression of the reporter gene was found to be greater in *C. glutamicum* transformants grown in acetate-containing medium than it was for transformants grown in glucose-containing medium (Wendisch *et al.*, *Arch Microbiol*. 168:262 (1997)) or other carbon sources (U.S. Patent No. 5,965,391).

[0012] Reinscheid et al., Microbiology 145:503 (1999), which is incorporated herein by reference in its entirey, discloses a transcriptional fusion between the C. glutamicum pta-ack promoter and a reporter gene (chloramphenicol acetyltransferase). C. glutamicum cells harboring the transcriptional fusion demonstrated enhanced reporter gene expression when grown in acetate-containing medium as compared to transformed cells that were grown in glucose-containing medium.

[0013] In P\u00e1tek et al., Microbiology 142:1297 (1996), which is incorporated herein by reference in its entirety, several DNA sequences from C. glutamicum, identified on the basis of their ability to promote the expression of a chloramphenicol resistance reporter gene in C. glutamicum cells, are disclosed and compared to one another in an attempt to define a consensus sequence for C. glutamicum promoters.

[0014] There is clearly a need for a broader assortment of well-defined Corynebacterium species promoters than has been heretofore described. Such promoters would be useful in the constitutive and/or regulated expression of genes in coryneform cells. For example, a collection of C. glutamicum promoters, regulated by inexpensive carbon sources, would facilitate the industrial-scale production of amino acids and purines in C. glutamicum cells by enhancing the expression of genes that encode components of the biosynthetic pathways for the desired amino acids or purines. Likewise, a versatile array of coryneform promoters would be useful for the industrial scale production of heterologous polypeptides in C. glutamicum cells by stimulating the enhanced expression of genes encoding such heterologous polypeptides.

BRIEF SUMMARY OF THE INVENTION

[0015] The present invention relates to isolated polynucleotides that function as transcriptional regulators, in particular, promoters, in Corynebacterium species host cells, preferably Corynebacterium glutamicum, Brevibacterium flavum, or Brevibacterium lactofermentum host cells, even more preferably C. glutamicum. host cells. These promoters are useful for regulating and enhancing the production of a variety of products in such host cells. Examples of products, the production of which may be enhanced in Corynebacterium species host cells as a result of the present invention, are amino acids, such as lysine; purine nucleotides, such as inosinic acid; and heterologous polypeptides. Since Corynebacterium species are especially useful for the industrial-scale production of amino acids, purines, and polypeptides, use of the promoters of the present invention may greatly improve the yields of these products from Corynebacterium species host cells.

[0016] In one embodiment of the present invention, an isolated polynucleotide is provided, comprising a first nucleic acid, the sequence of which is selected from the group consisting of SEQ ID NO:4 through 22.

[0017] Further embodiments of the invention include and isolated polynucleotide comprising a first nucleic acid, the sequence of which is at least 90% identical, and more preferably at least 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98% or 99% identical, to a sequence selected from the group consisting of SEQ ID NOs: 4 through 22.

[0018] Additional embodiments of the invention include an isolated polynucleotide comprising a first nucleic acid, the sequence of which comprises about 6 contiguous nucleotides, preferably about 10 contiguous nucleotides, even more preferably about 12, 15, 20, 30, 40, 50, 100, 150, 200, or 500 contiguous nucleotides, of a sequence selected from the group consisting of SEO ID NOs: 4

through 22, wherein the first nucleic acid is capable of regulating transcription of a second nucleic acid, preferably as part of a promoter.

[0019] In one aspect of this embodiment, the promoter comprises any one or more of the following genetic elements: a minus 10 ("-10") sequence; a minus 35 ("-35") sequence; a transcription initiation site; an enhancer region; and an operator region. Preferably the genetic elements are specific for Corynebacterium species, more preferably are specific for Corynebacterium glutamicum, Brevibacterium flavum, and Brevibacterium lactofermentum, and even more preferably are specific for Corynebacterium glutamicum.

[0020] Additional embodiments of the invention include an isolated polynucleotide comprising a first nucleic acid at least 10 nucleotides in length, preferably at least 12, 15, 30, 50 or 150 nucleotides in length, which hybridizes under stringent conditions to a reference nucleic acid, or the complement thereof, wherein the sequence of said reference nucleic acid is selected from the group consisting of SEQ ID NOs: 4 through 22. The meaning of the phrase "stringent conditions" as used herein is described infra.

[0021] In further embodiments of the present invention, an isolated polynucleotide is provided, comprising a first nucleic acid as described above; and a second nucleic acid. According to this embodiment, the first nucleic acid comprises a transcriptional regulatory region, preferably, a promoter.

[0022] In one aspect of this embodiment, said second nucleic acid encodes one or more polypeptides. Preferably, the physical location of the first nucleic acid relative to the second nucleic acid is such that, under the appropriate conditions, the first nucleic acid regulates transcription of the second nucleic acid, thereby facilitating production of the polypeptide.

[0023] In another aspect of this embodiment, the second nucleic acid encodes one or more components of a purine biosynthesis pathway. One example of a purine biosynthesis pathway included in this aspect of the invention is the enzymatic pathway that results in the synthesis of inosinic acid. [0024] In another aspect of this embodiment, the second nucleic acid encodes one or more heterologous polypeptides. The heterologous polypeptide may be one that is from a Corynebacterium species or one that is from a non-Corynebacterium species.

[0025] In one particularly preferred aspect of this embodiment, an isolated polynucleotide is provided, comprising a first nucleic acid as described above; and a second nucleic acid, which encodes a component of an amino acid biosynthesis pathway. Examples of amino acid biosynthesis pathways included in this aspect of the invention are the enzymatic pathways that result in the synthesis of L-glycine, L-alanine, L-leucine, L-methionine, L-phenylalanine, L-tryptophan, L-lysine, L-glutamine, L-glutamic acid, L-serine, L-proline, L-valine, L-isoleucine, L-cysteine, L-tyrosine, L-histidine, L-arginine, L-asparagine, L-aspartic acid, and L-threonine.

[0026] In a preferred aspect of this embodiment, the second nucleic acid encodes one of the following components of the L-lysine biosynthesis pathway: aspartokinase, aspartate beta-semialdehyde dehydrogenase, diaminopimelate dehydrogenase, diaminopimelate decarboxylase, dihydrodipicolinate synthetase, dihydrodipicolinate reductase, or pyruvate carboxylate.

[0027] Another embodiment of the invention provides an isolated polynucleotide comprising a first nucleic acid, the sequence of which is at least 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, or 99% identical to SEQ ID NO: 2, and a second nucleic acid which encodes a polypeptide which functions as a component of the lysine biosynthesis pathway. In this embodiment, the first nucleic acid regulates transcription of the second nucleic acid.

[0028] Yet another embodiment of the invention provides an isolated polynucleotide comprising a first nucleic acid, the sequence of which is at least 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, or 99% identical to SEQ ID NO: 3; and a second nucleic acid which encodes polypeptide which functions as a component of an amino acid biosynthesis pathway, preferably a lysine

biosynthesis pathway. In this embodiment, the first nucleic acid regulates transcription of the second nucleic acid.

Further embodiments of the invention provide isolated Corynebacterium [0029] chromosomes with a first nucleic acid integrated into the chromosome, the sequence of which is one of the following: a sequence which is at least 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, or 99% identical to SEQ ID NO:1;a sequence of which is at least 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%. 98%, or 99% identical to SEQ ID NO:23, a sequence of which is at least 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, or 99% identical to SEQ ID NO:30, or a sequence which is identical to either SEQ ID NO:26 or SEQ ID NO:27. In the embodiments where the integrated first nucleic acid has a sequence identical to either SEQ ID NO:26 or SEQ ID NO:27, the isolated chromosome also has a third nucleic acid integrated therein, the sequence of which is at least 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, or 99% identical to SEQ ID NO:28, operably linked to a transcription control region, wherein the nucleic acid encodes a polypeptide at least 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, or 99% identical to SEQ ID NO:29. In these embodiments, the isolated Corynebacterium species chromosome also has a second nucleic acid integrated therein, wherein said second nucleic acid encodes polypeptide which functions as a component of an amino acid biosynthesis pathway, preferably, a lysine biosynthesis pathway. Additionally in these embodiments, the first nucleic acid regulates transcription of the second nucleic acid.

[0030] In another aspect of the above embodiments, a Corynebacterium species host cell comprising a Corynebacterium species chromosome as described above is provided, preferably a host cell derived from Corynebacterium glutamicum, Brevibacterium flavum, and Brevibacterium lactofermentum. In a related aspect, a method of producing such a Corynebacterium species host cell is provided, which comprises transforming Corynebacterium species cells with a vector comprising a first nucleic acid as described above, wherein said vector facilitates

integration of the first nucleic acid into the chromosome of said Corynebacterium species cells, and selecting the host cell in certain preferred embodiments, the vector also comprises the second nucleic acid described above, and/or the third nucleic acid described above, with the first nucleic acid being physically situated to regulate transcription of the second nucleic acid. These vectors are also provided.

[0031] Examples of amino acid biosynthesis pathways which may be included in the above embodiments of the invention are the enzymatic pathways that result in the synthesis of L-glycine, L-alanine, L-leucine, L-methionine, Lphenylalanine, L-tryptophan, L-lysine, L-glutamine, L-glutamic acid, L-serine, Lproline, L-valine, L-isoleucine, L-cysteine, L-tyrosine, L-histidine, L-arginine, Lasparagine, L-aspartic acid, and L-threonine.

[0032] In those embodiments where the second nucleic acid encodes a component of the L-lysine biosynthesis pathway, the second nucleic acid may encode one or more of the following enzymes: aspartokinase, aspartate betasemialdehyde dehydrogenase, diaminopimelate dehydrogenase, diaminopimelate decarboxylase, dihydrodipicolinate synthetase, dihydrodipicolinate reductase, or pyruvate carboxylate.

[0033] In another embodiment of the present invention, a method is provided for producing a vector. More specifically, the method comprises inserting into a vector any of the isolated polynucleotides described herein.

[0034] In another embodiment of the present invention, a vector is provided comprising any of the isolated polynucleotides described herein.

[0035] In one aspect of this embodiment, the vector further comprises a multiple cloning region into which a heterologous second nucleic acid can be inserted, thereby allowing for the regulated and/or elevated expression of a variety of second nucleic acids therein

[0036] In another aspect of this embodiment, said vector is a shuttle vector. As used herein, the term "shuttle vector" refers to a vector that can replicate and be maintained in more than one host cell species. In a preferred aspect of this

embodiment, said shuttle vector can replicate and be maintained in a Corynebacterium species host cell, preferably a host cell derived from Corynebacterium glutamicum, Brevibacterium flavum, and Brevibacterium lactofermentum, and in an E. coli host cell.

[0037] In another embodiment of the present invention, a method of producing a transformed Corynebacterium species host cell is provided. The method of this embodiment comprises introducing into Corynebacterium species cells, preferably Corynebacterium glutamicum, Brevibacterium flavum, or Brevibacterium lactofermentum cells, a vector comprising any of the isolated polynucleotides described herein.

[0038] In one aspect of this embodiment, the polynucleotide, after being introduced into the host cell, is integrated into the chromosome of the host cell. In another aspect of this embodiment, the polynucleotide, after being introduced into the host cell, is maintained as an extrachromosomal element; i.e., it does not integrate into the chromosome of the host cell.

[0039] In another embodiment of the invention, a method is provided for the production of a biosynthetic product such as an amino acid, a purine nucleotide, or a heterologous polypeptide. According to this embodiment, a transformed Corynebacterium species host cell harboring a vector of the invention is used. More specifically, the vector comprises any isolated polynucleotide described herein.

[0040] In a preferred aspect of this embodiment, the polynucleotide comprises a first nucleic acid which regulates transcription of a second nucleic acid, where the second nucleic acid encodes a polypeptide which functions in an amino acid biosynthesis pathway, as described above. In this aspect, the first nucleic acid regulates transcription of the second nucleic acid, thereby resulting in elevated production of the amino acid by the transformed host cell.

[0041] In another aspect of this embodiment, the polynucleotide comprises a first nucleic acid which regulates transcription of a second nucleic acid, where the second nucleic acid encodes a component of a purine biosynthesis pathway. In this aspect, the first nucleic acid regulates transcription of the second nucleic acid, thereby resulting in elevated production of the purine by said transformed host cell.

[0042] In yet another aspect of this embodiment, the polynucleotide comprises a first nucleic acid which regulates transcription of a second nucleic acid, where the second nucleic acid encodes a heterologous polypeptide. In this aspect, the first nucleic acid regulates transcription of the second nucleic acid, thereby resulting in elevated production of the heterologous polypeptide by said transformed host cell.

[0043] In certain preferred aspects of this embodiment, the first nucleic acid comprises a promoter. Preferred promoters include, but are not limited to the following types of promoters: a constitutive promoter, an inducible promoter, a derepressable promoter, a heat sensitive promoter, and a cold sensitive promoter.

[0044] Where the promoter is an inducible promoter, the method for the production of a biosynthetic product as above may further comprise adding an inducer to the culture medium. The inducer may be present throughout the growth of the host cell, or alternatively, may be added to the culture medium after the host cell has grown to an optimal density. In a preferred aspect, the host cell is self-induced, i.e., the inducer is produced by said host cell at some point during its growth cycle. When the inducer is present in the culture medium, the first polynucleotide facilitates enhanced production of the biosynthetic product. Inducers which may be used in the invention include, but are not limited to acetic acid, pyruvate, ethanol, starch subunits, sugars, cellulose subunits, fatty acids, and triglycerides. Sugars that may be used as inducers in the present invention include, but are not limited to fructose, maltose, lactose, and arabinose.

Where the promoter is a derepressable promoter, the method for the production of a biosynthetic product as above may further comprise use of a culture medium which lacks a suppressor. In this aspect, the suppressor may be absent from the culture medium throughout the growth of said host cell, or the host cell may be initially cultured in a culture medium containing the suppressor.

[0045]

and be transferred to a culture medium lacking said suppressor when the host cell culture has reached an optimal density. In certain preferred aspects, the host cell may be self-regulating, i.e., the suppressor may be removed from said culture medium through metabolic depletion by the host cell. When the suppressor is removed or depleted from the culture medium, the first polynucleotide facilitates enhanced production of the biosynthetic product. Suppressors which may be used in the invention include, but are not limited to purines, pyrimidines, amino acids such as histidine, and oxygen.

[0046] Where the promoter is a heat sensitive promoter or a cold sensitive promoter, the method for the production of a biosynthetic product as above may comprise adjustment of the temperature of the growth medium to a temperature which is greater than or less than the optimal growth temperature for an untransformed *Corynebacterium* species cell. For example, the yield of the biosynthetic product may be increased when said host cell is grown at a temperature adjusted to be greater than the optimal growth temperature for an untransformed *Corynebacterium* species cell, or the yield of the biosynthetic product may be increased when the host cell is grown at a temperature adjusted to be less than the optimal growth temperature for an untransformed *Corynebacterium* species cell.

DETAILED DESCRIPTION OF THE INVENTION

[0047] The present invention relates to isolated polynucleotides that function as transcriptional regulators, in particular, promoters, in Corynebacterium species host cells, preferably Corynebacterium glutamicum, Brevibacterium flavum, or Brevibacterium lactofermentum host cells, even more preferably C. glutamicum. host cells. These promoters are useful for regulating and enhancing the production of a variety of products in such host cells. Sources of promoters include nucleotide sequences from the 5' end of native chromosomal genes from Corynebacterium species, from sequences on plasmids that replicate in Corynebacterium species, from sequences in the genome of phage that infect Corynebacterium species, from sequences derived from other microorganisms, e.g., Escherichia coli, or from sequences assembled by humans (tac, trc) which are not found in nature. Genes of ribosomal proteins, ribosomal RNAs and elongation factors show high levels of expression. The promoters of these genes are candidates for increasing expression of amino acid biosynthetic pathway genes.

[0048]

Another reason for changing promoters of genes in biosynthetic pathways is to make the pathway independent of factors that control the pathway in the wild type organism. For example the native promoter of the operon that contains diaminopimelate decarboxylase of the lysine biosynthetic pathway of *C. glutamicum* can respond to arginine or lysine in the growth medium. Arginine increased transcription three-fold and lysine decreased transcription by one third (Oguiza, et al., J Bact. 175:7356-7362 (1993)). Diaminopimelate decarboxylase activity decreased 60% in cells grown in minimal medium supplemented with 10mmM lysine (Cremer, et al., J Gen Microbiol. 134:3221-3229 (1988)). Replacing the promoter of lysA which encodes the diaminopimelate decarboxylase is one way to make lysine biosynthesis independent of arginine and lysine levels in media.

[0049]

Gene expression from many of the transcriptional regulators of the present invention can be easily and inexpensively regulated by manipulating the composition and/or temperature of the medium in which cells containing these promoters are cultured. Regulated expression from a promoter is especially useful for controlling the production of a gene product that is toxic to cells at high levels. To allow for maximum production of a gene product, a culture of cells possessing the gene of interest under the control of an inducible promoter can be

first grown to a sufficient cell density under conditions in which expression from the promoter is suppressed. Then, by manipulating the culture medium, expression from the promoter is induced. This strategy assures not only a high level of gene expression in the individual cells, but also a sufficiently high number of cells in the culture, so that maximum production of the gene product by the entire cell population is achieved.

A. Definitions

[0050] In order to provide a clear and consistent understanding of the specification and claims, including the scope to be given such terms, the following definitions are provided. It is also to be noted that the term "a" or "an" entity, refers to one or more of that entity; for example, "a polynucleotide," is understood to represent one or more polynucleotides. As such, the terms "a" (or "an"), "one or more," and "at least one" can be used interchangeably herein.

[0051] Auxotroph. As used herein, the term refers to a strain of microorganism requiring for growth an external source of a specific metabolite that cannot be synthesized because of an acquired genetic defect.

[0052] Amino Acid Supplement. As used herein, the term refers to an amino acid required for growth and added to minimal media to support auxotroph growth.

[0053] Chromosomal Integration. As used herein, the term refers to the insertion of an exogenous DNA fragment into the chromosome of a host organism; more particularly, the term is used to refer to homologous recombination between an exogenous DNA fragment and the appropriate region of the host cell chromosome

[0054] Corynebacterium species. As will be understood by those skilled in the art, the terms "Corynebacterium" or "Corynebacterium species" includes those organisms previously identified in the literature as "Brevibacterium species," for example Brevibacterium flavum and Brevibacterium lactofermentum which have

now been reclassified into the genus Corynebacterium. (Int. J. Syst. Bacteriol. 41: 255 (1981)). Accordingly, the term "Corynebacterium species" is used herein interchangably with "Corynebacterium species and Brevibacterium species."

[0055] Enhancers. As used herein, the term refers to a DNA sequence which can stimulate promoter activity and may be an endogenous element or a heterologous element inserted to enhance the level, i.e., strength of a promoter.

[0056] Inducer. As used herein, the term "inducer" refers to molecule which acts to stimulate transcription from an inducible promoter. The inducer may be produced by a host cell, or added to a culture medium in which the host cell is being grown.

[0057] Isolated Polynucleotide. As used herein, the term is intended to mean a polynucleotide, DNA or RNA, which has been removed from its native environment. For example, recombinant DNA molecules contained in a vector are considered isolated for the purposes of the present invention. Further examples of isolated DNA molecules include recombinant DNA molecules maintained in heterologous host cells or purified (partially or substantially) DNA molecules in solution. However, a nucleic acid molecule contained in a clone that is a member of a mixed clone library (e.g., a genomic or cDNA library) and that has not been isolated from other clones of the library (e.g., in the form of a homogeneous solution containing the clone without other members of the library) or a chromosome isolated or removed from a cell or a cell lysate, is not "isolated" for the purposes of this invention. Isolated RNA molecules include in vivo or in vitro RNA transcripts of the DNA molecules of the present invention. Isolated nucleic acid molecules according to the present invention further include such molecules produced synthetically.

[0058] Lysine Biosynthetic Pathway Protein. As used herein, the term "lysine biosynthetic pathway protein" is meant to include those peptides, polypeptides or proteins, and enzymes, which are directly involved in the synthesis of lysine from aspartate. These proteins may be identical to those which naturally occur within a host cell and are involved in the synthesis of lysine within that host cell.

Alternatively, there may be modifications or mutations of such proteins, for example, the proteins may contain modifications or mutations which do not significantly affect the biological activity of the protein. For example, the natural protein may be modified by mutagenesis or by introducing or substituting one or more amino acids, preferably by conservative amino acid substitution, or by removing nonessential regions of the protein. Such modifications are readily performed by standard techniques. Alternatively, lysine biosynthetic proteins may be heterologous to the particular host cell. Such proteins may be from any organism having genes encoding proteins having the same, or similar, biosynthetic roles.

[0059]

Mutagenesis. As used herein, the term refers to a process whereby a mutation is generated in DNA. With "random" mutagenesis, the exact site of mutation is not predictable, occurring anywhere in the genome of the microorganism, and the mutation is brought about as a result of physical damage caused by agents such as radiation or chemical treatment. rDNA mutagenesis is directed to a cloned DNA of interest, and it may be random or site-directed.

[0060]

Mutation. As used herein, the term refers to a one or more base pair changes, insertion or deletion in the nucleotide sequence of interest.

[0061]

Operably Associated. As used herein, the term "operably associated" refers to a association of nucleic acid elements in a functional relationship. A nucleic acid is "operably associated" when it is placed into a functional relationship with another nucleic acid sequence. For instance, a promoter or enhancer is operably associated with a polypeptide coding region if it affects the transcription of the polypeptide coding region. Operably associated means that the nucleic acids being associated are typically close together or contiguous and, where necessary, join two polypeptide coding regions, contiguous and in reading frame. However, since enhancers generally function when separated from the promoter by several kilobases, some polynucleotide elements may be operably associated but not close together or contiguous.

[0065]

[0062] Operon. As used herein, the term refers to a contiguous portion of a transcriptional complex in which two or more open reading frames encoding polypeptides are transcribed as a multi-cistronic messenger RNA, controlled by a cis-acting promoter and other cis-acting sequences necessary for efficient transcription, as well as additional cis acting sequences important for efficient transcription and translation (e.g., mRNA stability controlling regions and transcription termination regions).

[0063] Parental Strain. As used herein, the term refers to a strain of host cell subjected to some form of treatment to yield the host cell of the invention.

[0064] Phenotype. As used herein, the term refers to observable physical characteristics dependent upon the genetic constitution of a host cell.

Promoter. As used herein, the term "promoter" has its art-recognized meaning, denoting a portion of a gene containing DNA sequences that provide for the binding of RNA polymerase and initiation of transcription and thus refers to a DNA sequence capable of controlling the expression of a coding sequence or functional RNA. Promoter sequences are commonly, but not always, found in the 5' non-coding regions of genes, upstream of one or more open reading frames encoding polypeptides. Sequence elements within promoters that function in the initiation of transcription are often characterized by consensus nucleotide sequences. The promoter sequence includes proximal and more distal upstream elements. Examples of proximal elements in bacterial promoters include a -10 region and a -35 region, which are discussed in more detail, infra. Examples of more distal elements include operator regions and enhancer regions. As used herein, the term "endogenous promoter" refers to a promoter sequence which is a naturally occurring promoter sequence in that host microorganism. The term "heterologous promoter" refers to a promoter sequence which is a non-naturally occurring promoter sequence in that host microorganism. The non-naturally occurring promoter sequence may be from any prokaryotic or eukaryotic organism. A synthetic promoter is a nucleotide sequence, having promoter activity, and not found naturally occurring in nature.

[0066]

Promoters may be derived in their entirety from a native gene, or be hybrid promoters. Hybrid promoters are composed of different elements derived from different promoters found in nature, or even comprise synthetic DNA segments. Hybrid promoters may be constitutive, inducible or environmentally responsive.

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Useful promoters include constitutive and inducible promoters. Many such promoter sequences are known in the art. See, for example, U.S. Pat. Nos. 4,980,285; 5,631,150; 5,707,828; 5,759,828; 5,888,783; 5,919, 670, and, Sambrook, et al., Molecular Cloning: A Laboratory Manual, 2nd Ed., Cold Spring Harbor Press (1989). Other useful promoters include promoters which are neither constitutive nor responsive to a specific (or known) inducer molecule. Such promoters may include those that respond to developmental cues (such as growth phase of the culture), or environmental cues (such as pH, osmoticum, heat, dissolved gases, or cell density).

100681

Examples of environmental conditions that may effect transcription by inducible promoters include anaerobic conditions, elevated temperature, reduced temperature, or the presence of light. It is understood by those skilled in the art that different promoters may direct the expression of a gene in different cell types, or in response to different environmental conditions. Promoters which cause a gene to be expressed in most cell types at most times are commonly referred to as "constitutive promoters". It is further recognized that since in most cases the exact boundaries of regulatory sequences have not been completely defined, DNA fragments of different lengths may have identical or similar promoter activity.

[0069]

Relative Growth. As used herein, the term refers to a measurement providing an assessment of growth by directly comparing growth of a parental strain with that of a progeny strain over a defined time period and with a defined medium.

[0070]

Stringent Hybridization Conditions. As used herein, the term "stringent hybridization conditions" is intended to mean overnight incubation at 42 °C in a solution comprising: 50% formamide, 5x SSC (750 mM NaCl, 75mM trisodium

citrate), 50 mM sodium phosphate (pH 7.6), 5x Denhardt's solution, 10% dextran sulfate, and 20 g/ml denatured, sheared salmon sperm DNA, followed by washing the filters in 0.1x SSC at about 65°C.

[0071] Suppressor or Repressor. As used herein, the terms "suppressor" or
"repressor" refer to molecules which act to block or reduce transcription from an
derepressable promoter. The suppressor or repressor may be produced by a host
cell, or added to a culture medium in which the host cell is being grown.
Furthermore, during the growth of a host cell, the suppressor substance can be
metabolized by the host cell, thereby removing it from the culture medium,
thereby increasing transcription from the derepressable promoter.

[0072] Transcription factor. As used herein, the term "transcription factor" refers to RNA polymerases, and other proteins that interact with DNA in a sequence-specific manner and exert transcriptional regulatory effects. Transcriptional factors may be transcription inhibitory proteins or transcription activator proteins. In the context of the present invention, binding sites for transcription factors (or transcription complexes) are often included in the transcriptional regulatory element(s).

[0073] Transcription factor recognition site. As used herein, a "transcription factor recognition site" and a "transcription factor binding site" refer to a polynucleotide sequence(s) or sequence motif(s) which are identified as being sites for the sequence-specific interaction of one or more transcription factors, frequently taking the form of direct protein-DNA binding. Transcription factor recognitions sites which bind transcription inhibitory proteins or transcription activator proteins are also referred to as "operator regions." Typically, transcription factor binding sites can be identified by DNA footprinting, gel mobility shift assays, and the like, and/or can be predicted on the basis of known consensus sequence motifs, or by other methods known to those of skill in the art.

[0074] Transcriptional Complex. As used herein, the term "transcriptional unit" or "transcriptional complex" refers to a polynucleotide that comprises one or more coding regions for polypeptides, a cis-acting linked promoter and other

cis-acting sequences necessary for efficient transcription of the structural sequences, distal regulatory elements necessary for appropriate transcription of the structural sequences, and additional cis sequences important for efficient transcription and translation (e.g., mRNA stability controlling regions and transcription termination regions). In bacteria, a transcriptional complex may comprise a single coding region, or one or more coding regions, e.g., as part of an operon.

[0075] Transcriptional Regulatory Element. As used herein, the term "transcriptional regulatory element" refers to a DNA region which activates transcription alone or in combination with one or more other DNA regions. A transcriptional regulatory element can, for example, comprise a promoter, response element, negative regulatory element, silencer element, gene suppressor, transcription terminator, and/or enhancer.

B. Microbiological and Recombinant DNA Methodologies

[0076] The invention as provided herein utilizes some methods and techniques that are known to those skilled in the arts of microbiology and recombinant DNA technologies. Methods and techniques for the growth of bacterial cells, the introduction of isolated DNA molecules into host cells, and the isolation, cloning and sequencing of isolated nucleic acid molecules, etc., are a few examples of such methods and techniques. These methods and techniques are described in many standard laboratory manuals, such as Davis et al., Basic Methods In Molecular Biology (1986), J.H. Miller, Experiments in Molecular Genetics, Cold Spring Harbor Laboratory Press, Cold Spring Harbor, New York (1972); J.H. Miller, A Short Course in Bacterial Genetics, Cold Spring Harbor Laboratory Press, Cold Spring Harbor, New York (1992); M. Singer and P. Berg, Genes & Genomes, University Science Books, Mill Valley, California (1991); J. Sambrook, E.F. Fritsch and T. Maniatis, Molecular Cloning: A Laboratory Manual, 2d ed., Cold Spring Harbor, New

York (1989); P.B. Kaufman et al., Handbook of Molecular and Cellular Methods in Biology and Medicine, CRC Press, Boca Raton, Florida (1995); Methods in Plant Molecular Biology and Biotechnology, B.R. Glick and J.E. Thompson, eds., CRC Press, Boca Raton, Florida (1993); and P.F. Smith-Keary, Molecular Genetics of Escherichia coli, The Guilford Press, New York, NY (1989), all of which are incorporated herein by reference in their entireties.

[0077] Unless otherwise indicated, all nucleotide sequences newly described herein were determined using an automated DNA sequencer (such as the Model 373 from Applied Biosystems, Inc.). Therefore, as is known in the art, for any DNA sequence determined by this automated approach, any nucleotide sequence determined herein may contain some errors. Nucleotide sequences determined by automation are typically at least about 90% identical, more typically at least about 95% to at least about 99.9% identical to the actual nucleotide sequence of the sequenced DNA molecule. The actual sequence can be more precisely determined by other approaches including manual DNA sequencing methods well known in the art.

C. Polynucleotides

[0078] Certain embodiments of the present invention are directed to an isolated polynucleotide comprising a first nucleic acid, the sequence of which is related to, or identical to, a nucleotide sequence, or fragment thereof, selected from the group consisting of SEQ ID NO:1, SEQ ID NO:2, SEQ ID NO:3, SEQ ID NO:4, SEQ ID NO:5, SEQ ID NO:6, SEQ ID NO:7, SEQ ID NO:8, SEQ ID NO:9, SEQ ID NO:10, SEQ ID NO:11, SEQ ID NO:12, SEQ ID NO:13, SEQ ID NO:14, SEQ ID NO:15, SEQ ID NO:16, SEQ ID NO:17, SEQ ID NO:18, SEQ ID NO:19, SEQ ID NO:20, SEQ ID NO:21, SEQ ID NO:22, SEQ ID NO:23, SEQ ID NO:24, SEQ ID NO:26, SEQ ID NO:27, SEQ ID NO:28, and SEQ ID NO:30. SEQ ID NO:1-3 have been previously described. See, e.g., Reinscheid et al., Microbiology 145:503 (1999); U.S. Patent No. 5,700.661; and U.S. Patent No.

5,965,391. SEQ ID NOs 23, 24, 26, 27, 28, and 30 have been previously described. See, e.g., Ben-Samoun et al., FEMS Microbiology Letters 174:125-130 (1999), Brosius, J. et al., J. Biol. Chem. 260, 3539-3541 (1985), Amann, E., et al., Gene 69: 301-315 (1988), Moeckel, et al., J.Bacteriol. 174:8065-8072 (1992), Keilhauer, et al. J.Bacteriol 175:5595-5603 (1993) and Patek et al., Appl. Env. Microbiol., 60:133-140 (1994). Isolation of polynucleotides comprising nucleic acids the sequence of which are identical to SEQ ID NOs 4-22 are described herein. While not being bound by theory, it is believed that polynucleotides having these sequences can be used to regulate gene expression in Corynebacterium species. The genes regulated by SEQ ID NOs 4-22, determined by similarity of the genes to genes identified in other organisms, are listed in Table 1A, along with putative exogenous regulatory molecules. SEQ ID NOs 1-22 are listed in Table 1B.

TABLE 1A

Nucleotide sequences that can be used to regulate gene expression

Seq. I.D. NO:	Gene*	Regulatory Molecule [‡]
1	pta	acetate
2	aceA	acetate
3	aceB	acetate
4	adh	ethanol
5	aldB	ethanol
6	poxB	pyruvate
7	ldh	pyruvate
8	amyE	carbon
9	malZ	carbon
10	bglX	carbon
11	gam	carbon
12	glgX	carbon
13	hisD	histidine
14	pyrR	pyrimidine
15	purD	purine
16	hrcA	temperature
17	htpX	temperature

18	dnaK	temperature
19	ctc	temperature
20	grpE	temperature
21	clpB	temperature
22	narA	oxygen

Sequence I.D. NOs 1, 2, and 3 have been previously described. The remaining sequences were discovered in ADM's Corynebacterium glutamicum genome sequencing project.

^{*} Putative genes regulated by sequence I.D. NOs 4-22 were determined by homology to genes identified in other organisms, e.g., Escherichia coli or Bacillus subtilis.

[‡] Putative regulatory molecules associated with the regulatory regions of SEQ I.D. NOs 4-22 were determined by analogy to regulatory regions identified in other organisms.

Table 1R

Nucleotide sequences that can be used to regulate gene expression

SEQ ID NO: 1

SEQ ID NO:2

SEQ ID NO:3

SEQ ID NO:4

SEO ID NO:5

CGAT CATCGACTTGGCGGAGARAACCGGAACCTCAAAGATCCCTGGCTAAGGTCACACTGGGAATGA CACACTCAAAAGTCAGTTACCACCACGCGGCTCCCCCCCGCGGCGGCTGGTCACCGCGCCGCCCCCCCGCG GTCGTCATCTGGGAACGCCGCAAGGGGCGTTCCAAGTAAAAACATCCTTGTCACGCCCGCTTCTACCAC ATTAAGCGGGCACCTCCATTTACTTTTGGAGGTGCCCCTTTCGCCTTTGGCAATTAGATTACTGCAT AACCACCCGAACAGGGGTAATAACTTTTGAAAGCCTTTCGCCCTTCAGCTACGAATTAGATTACAGCAT GGCGTGAATTTAACAGGGGTTCTTACCCCGCATTTAGATTTCGTTTTGGTAATTAGGTCTCCTAAACCTGCCGAAT TGCGCGTTATTCACAATCGTTGACCAAGTGCCACCTGACGCACAGGTAGTGCTCAGGGGTGCCCCAA GGCGGCATATTCACAATCGTTGACCAAGTGCCACCTGACGCACAGGTAGTGCTCAGGGTGGCCCCAA GGCGGCATTTCACAATCGTTGACCAAGTGCCACCTGACGCACAGGTAGTGCTCAGGGTGGGCCCCAA

SEQ ID NO:6

TTTTAGACCACGGGGCTGTGTGGGGATTTAAGACCTCGGAAATGCTAGGGGACTGCAGTGTGAGTGGG TTCTTTGAGGCGCTTAGAGGCGATTCTGTGAGGCTGACTATTTTGGGGCTGGGGTGAAATTGGCAGT TTTGAGGCGACCAACAGGCGTACAACATGACTAAAAAACCAAGTTTTGGGAGGTGGTTATAGTTGGCAGTGCTTGGTTT CTGGGGCGCGACCAGCAGCTGGAGAAGTTGGGCAATTGGCAGTTTTGGCAGCTGTCTGGT TTTGAGGGCCGAACTGCCATGAACTAGTGGCAATCGCAACAGCTTTTGGCCTCCAGCGCTGT TTTGAGGCCCAACTGCAATTGAACAACAGCAGCTAGTAGGCAATTGGCATCGCAGCAGCTGT TTTGAGCACCAACCCCCTTAAACAACAGCAGTAGGAAACAGCAGTAGAGAAAATTG

SEQ ID NO:7

SEO ID NO:8

SEO ID NO:9

GGGCATCATGGCCATTGTTGGTGGATTCACCATGCCTGTCGCAATCGCTAGGACCAGGGATAAGAACCTC GTGTGGTTCCCCGTGGTCTTTGGTGCATCGATGTTCCTCGGTTATGTGGGAACGTGGCTGTGGCCGTCCC AAGGCTGGTACCTGTGGTCATTCCTTCTTGGTTTAGGTGGACTCTGCTTCCCGATGGCTATCGCCCTGAT TCCAGCGCGTACGAAAGATCCGAGAATTACCGCAAGCTTGTCTGGATTTGTGCAGCCGGTGGGTTACATT CTTGCAGCCCTTGGGCCATTGGCAGTGGGAGCGATCTACCAGGCGATTGGCTCCTGGTCAGAGATCCTCG TTGGTTTGGCCTTGGGCACAATAGTGTTGTCGATTGTGGGATTCAGAGCAGCACGCAATGTGACGGTTGA TGATGAATTGAGGAGATCAAAGTAGCCTCAACTAAGCGTCGCGATAAGAACGAGGGCAAGGCTGATGTA CTCTGTCAACCATGGATAAACCGGTCGTGAGGGATGCAGCTCTGCTGATTTTTCGCGCTGTGCTCGGAGT GATCTTTGTGGCACACGGGTGGGAAAAGCTGTTCATCTCCGGAGTTACCAAGACAACAGGACAATTTTCA GCCTGGGGAGTGCCTCAACCCAAGCTCTCGGTGTGGATCACATCGATCTCTGAGCTGCTCGGTGGTGCCT TCCTAGTGGTTGGTTGCTCACCACCTTTGTTGCTGGTGCATTAGCGCTGTTGATCGCCGCTGCTATTTA CTTTGTGCACTTGAGTTCGGGCTTTTTCACAGTTGATAACGGCATCGAATTCCCCTTGCTCATCATTGTT TCTTTGCTCGTGATCGTTGTTTTGGTTCTGGTAGAGCCAGCGTTGATGGGGTGCTCACGCGTGGTTGAC TGTAGTGCCATTCAAGCCGCGCTGTCCGCCAAATTAGATGGTGAGCCGACAGGCTTGGATGATGCAGTAA TTGAAGCGCACCTCGCTAATTGTGAAGAGTGCAGAAATTACTACAACCGTGCTGCTGAGTTGAATCGGAT GCTCAATTTTTGCGCCGCGGAACCTCGCACCCTGACCCCGCCTGATCTATCAGAGATCATTCTGGCAGAG GTGGAACCAGAATGGCGCAGGCATGCCAACGCCAAGGTTGTGGGATCCCTGCTATCGCGAGTGTTGTTGG TGATCCTGGGTGTGGTTTACCTCGCCTGGGGTATCACAATGTTGGGGGATTCGGCGTCGATAAGCGTCCA AGAAGACCCGCTCACCTCGCGCCTGCTCGCGGAGGCCGTTGCCTACCGCATTGCTCTTTCTGTGGGGCTG TTATTTGCGGCGTGGAAGCCGCGGATTATCGCGGGCATGCTCCCGATTTTTGGAACGTTGTGGACATTTA GTGCTGGTTTTGCTGCGCGCGATCTCGTGTTTGGCGTCGCCGATTCACAGACGGGACTGTCCATTGGTCT GCTGTTGATTTCTACGATTGTGCTGTCGTTGCGTTGGTGAATAGTTCTGGACCGGGTATTTTGCGGCCG ACATGGAACTCATTGAACGCCGCGCCCGGCTAAGGTGGGAGGC

SEO ID NO:10

TABATA-CGGCATCCGGCCTGAAAGGACAGTGTGGAACCCAAAAATCATGCCCTGGCAGGAATCGTGTTTAAG GGGTTAAAACGCATCGACCCCATCCAGCCCATCCGGCAACTGACTCGACTGCTGATAAAGCGATTCTGGTTG GTGASGTGTGAAAACGGACTGCGGGGGGCCAATTCACCAAAAATCCATACGAGTGTGCCACCCGATTTTC ACATGCGTTGGAGACCTCCCTTTTTTGACCACTTTAATTGCTGAACCCCGTATAGGGAGAAGGGGTACCAACCGATTTTC ACATGCGTTGCTTTTTTTTGGGGGGAATCTGACTACCATGGTAAGAAACAGGGAAAGGGGTACATTTTGGACA CAAGACTAGCTCAAATCGGCTCCGCCTCTTCTGTCTATTGACAGAGGGCCTCACATTTCGCCGATC TCGACGGCGACGGCGTACTTGCACCTTATGAAGATTGGCTCTAACCCCCAGCAGAGCGTCCGCTCACTC

SEQ ID NO:11

SEQ ID NO:12

GGTGTCAGTGTATCCGCGAACCAGAGCCTCGATGGCAGCTGTGTCTTCTGGGGAAGCGGTCTGGCCGTTT TCAACTGGCGCAGCTTCAGCGAAGGTAATTGGGTTCTCAGACAGCTCTTGTGACAGTACATCAAGCTGTG CTTGCTGCTCTTCGTTGATGGTTGTTGCTTCGGAAGATTCAGATTCAGATTCAGATTCGGACGAGGTGGT TTCTGCTGCTTCCGCCGAGGTCGTTGCGCTGGAAGAGGAAGAATTTGTCGTCGTCGCTGCGCTGGATGAG GCTGCTTCTGTATCAGATGACTCACTGCTACATGCGGTTAAGAGAAGTGGGGTGACCATGAGTGCTGCGA AAGCAGCCTTCTTTGAGGAAAGGCGAATAGACAAAGTTCTGCTCCTGATAAATCATCGACATGCTCCGGA AAACTTAAAAATTCCCGGACGGTTCACGCAGATTACCCTAGCAAAGCAATCTAGCTGACGACCCAATTTA GTCCTGTCATTATGCTGGCAATTGTGCAGCTATCTAAAGAATCTATTATTGGGGCAGCCGTTTCGATCCT GAGCGAGTTCGGTTTGTCGGATATGACCATGCGTCGCGTCGCAAAGCAATTAAATGTCGCGCCGGGCGCG CTGTATTGGCATTTTAAAAATAAGCAGGAGCTTATCGACGCCACCTCACGCCATCTCCTGGCGCCTATCT TGGGGCGCAACGACGAGCAGCAAGCATTTCCGCGCAGGAAACGTGCGCAGAAATGCGTTCACTGAT GATGCAAACCAAAGACGGTGCGGAAGTCATCAGTGCCGCACTGAGTAATCAGCAACTGCGCCAAGAATTG GAATCTCTCATTTCCGACTCTTTAAAAGAACCTAATGAGGTCGGTGCTTTTACGCTGCTGCATTTTGTGG TGGGTGCAGTATTAACAGAACAAACTCAGCTGCAGATGCACGAGTTCACGGCTGGCGCGGAAGATGACAC CTAGACGCGCTTGGGCATATAAGATAGCGTTCT

SEO ID NO:13

SEO ID NO:14

SEQ ID NO:15

SEO ID NO:16

SEQ ID NO:17

SEQ ID NO:18

AGACTTGGCGTCAAAGTTGAGTGGGACCTGTTGAAAACAGCTCGTATCTTCCCTGATTGGGTGGTTGAAA TTAGGGGTAAACCGGATTTTTCTCAAGTGAAGGGCTTTGACCTGTGTAAATTAAGAAAGTTGAGTCAT GTGGGACAACTTTGTGGGCATTTTTACCTTTGCCTTATGTTAAAGTTGAGTTGAGCAGCGCGAGCCAATTAGGAAG TTTTTCTTACCGGCGAAAGTCGGTGGAAGCAAGTCAAAGCTCAAGCCGTGGACAATACTAAAATCACCTA AAACAGGAGGACACATT

SEQ ID NO:19

SEQ ID NO:20

SEQ ID NO:21

SEQ ID NO:22

TGGCCAPTCCTGTGGTCCATGCTCGCCCTGTTCTTCTCCTGGACTGGGCAACGCCGGCACATTCAAAC
AAATGCCCATACATTTGCCCAAACGCCAAGCAGTGGCGTGATTCGAAC
TCGGCCCCTTCATTGTCGGTGCTCTCCTCCTCCATCCACTGTGCGTTCCTTCTGGGGCTGCGTTGTGTGCTT
CGGCCCCTTCATTGCCTGGGTTTGCCTTGGATTCTCTCACTGTTTCTGGGGCTGCTGTGTG
TTCTTCTACATGGCACCGGCTTTGACCTGGATCTTCATCTACTGGAACGTCTTCTGAACAGTCTTCTGAAAAGGTCA
CCGAAATCCAATGACTACATCATCTTCTTCTGGGAAGTCTTCTGAACAGTCTTCTGAAAAGATCA
ACCCCCTCTTCAAACGTCGGACTTTCCTAAAGAAAAGGCACGCGAAAGGCCAGCAGAATTTTCCT
TCAGGGGGAGCCCAAGCCGATGTTTCTAAAAAAAGCACCGAATGGCGTTCCGAAAAAGCACCGCATTTTCTCAT
CACACATGGC
CACACATGGC

[0079]

In certain embodiments, polynucleotides of the invention comprise a first nucleic acid, the sequence of which is at least 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98% or 99% identical to a sequence selected from the group consisting of SEQ ID NO:4, SEQ ID NO:5, SEQ ID NO:6, SEQ ID NO:7, SEQ ID NO:8, SEQ ID NO:9, SEQ ID NO:10, SEQ ID NO:11, SEQ ID NO:12, SEQ ID NO:13, SEQ ID NO:14, SEQ ID NO:15, SEQ ID NO:16, SEQ ID NO:17, SEQ ID NO:18, SEQ ID NO:19, SEQ ID NO:20, SEQ ID NO:21, and SEQ ID NO:22, or a complementary sequence thereof. Of course, this embodiment also encompasses a first nucleic acid, the sequence of which is identical to any of the recited nucleic acid sequences.

[0800]

By a polynucleotide comprising a first nucleic acid, the sequence of which is at least, for example, 95% "identical" to a reference nucleotide sequence is intended that the first nucleic acid sequence is identical to the reference sequence except that the first nucleic acid sequence may include up to five mismatches per each 100 nucleotides of the reference nucleic acid sequence. In other words, to obtain a first nucleic acid, the sequence of which is at least 95% identical to a reference nucleic acid sequence, up to 5% of the nucleotides in the reference sequence may be deleted or substituted with another nucleotide, or a number of nucleotides up to 5% of the total nucleotides in the reference sequence may be

inserted into the reference sequence. The reference (query) sequence may be any one of the entire nucleotide sequences shown in SEQ ID NO:4, SEQ ID NO:5, SEQ ID NO:6, SEQ ID NO:7, SEQ ID NO:8, SEQ ID NO:9, SEQ ID NO:10, SEQ ID NO:11, SEQ ID NO:12, SEQ ID NO:13, SEQ ID NO:14, SEQ ID NO:15, SEQ ID NO:16, SEQ ID NO:17, SEQ ID NO:18, SEQ ID NO:19, SEQ ID NO:20, SEQ ID NO:21, or SEQ ID NO:22, or any fragment of any of these sequences, as described *infra*.

[0081]

As a practical matter, whether any particular nucleic acid sequence is at least 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98% or 99% identical to, for instance, a nucleotide sequence selected from the group consisting of SEQ ID NO:4, SEQ ID NO:5, SEQ ID NO:6, SEQ ID NO:7, SEQ ID NO:8, SEQ ID NO:9, SEO ID NO:10, SEQ ID NO:11, SEQ ID NO:12, SEQ ID NO:13, SEQ ID NO:14, SEQ ID NO:15, SEQ ID NO:16, SEQ ID NO:17, SEQ ID NO:18, SEO ID NO:19, SEQ ID NO:20, SEQ ID NO:21, and SEQ ID NO:22, or a complementary sequence thereof, can be determined conventionally using sequence analysis computer programs such as a OMIGA® Version 2.0 for Windows, available from Oxford Molecular, Ltd. (Oxford, U.K.). OMIGA uses the CLUSTAL W alignment algorithm using the slow full dynamic programming alignment method with default parameters of an open gap penalty of 10 and an extend gap penalty of 5.0, to find the best alignment between two nucleotide sequences. When using CLUSTAL W or any other sequence alignment program to determine whether a particular sequence is, for instance, 95% identical to a reference sequence according to the present invention, the parameters are set, of course, such that the percentage of identity is calculated over the full length of the reference nucleotide sequence such that gaps, mismatches, or insertions of up to 5% of the total number of nucleotides in the reference sequence are allowed.

[0082]

This embodiment of the present invention is directed to polynucleotides comprising a first nucleic acid, the sequence of which is at least 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98% or 99% identical to a nucleic acid sequence selected from the group consisting of SEQ ID NO:4, SEQ ID NO:5, SEQ ID

NO:6, SEQ ID NO:7, SEQ ID NO:8, SEQ ID NO:9, SEQ ID NO:10, SEQ ID NO:11, SEQ ID NO:11, SEQ ID NO:12, SEQ ID NO:13, SEQ ID NO:14, SEQ ID NO:15, SEQ ID NO:16, SEQ ID NO:17, SEQ ID NO:18, SEQ ID NO:19, SEQ ID NO:20, SEQ ID NO:21, and SEQ ID NO:22, or a complementary sequence thereof, irrespective of whether they have functional activity. This is because even where a particular polynucleotide does not have functional activity, one of skill in the art would still know how to use the nucleic acid molecule, for instance, as a hybridization probe, an S1 nuclease mapping probe, or a polymerase chain reaction (PCR) primer.

100831

Preferred, however, are polynucleotides comprising a nucleic acid, the sequence of which is at least 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98% or 99% identical to a nucleic acid sequence selected from the group consisting of SEQ ID NO:4, SEQ ID NO:5, SEQ ID NO:6, SEQ ID NO:7, SEQ ID NO:8, SEQ ID NO:9, SEO ID NO:10, SEO ID NO:11, SEQ ID NO:12, SEQ ID NO:13, SEQ ID NO:14, SEQ ID NO:15, SEQ ID NO:16, SEQ ID NO:17, SEQ ID NO:18, SEO ID NO:19, SEO ID NO:20, SEQ ID NO:21, and SEQ ID NO:22, or a complementary sequence thereof, which do, in fact, have functional activity, in particular, polynucleotides which are capable of regulating transcription in Corvnebacterium species. Examples of polynucleotides which regulate transcription include promoters, operators, transcription factor recognitions sites, transcriptional complexes, and transcriptional regulatory elements, as described herein. Preferably, polynucleotides of the present invention comprise a promoter which functions in Corvnebacterium species, preferably in Corvnebacterium glutamicum. Brevibacterium flavum, or Brevibacterium lactofermentum, even more preferably in C. glutamicum. Assays to determine whether a polynucleotide is capable of regulating transcription in Corynebacterium species can be routinely performed using techniques described herein and otherwise known in the art.

[0084]

Accordingly, the present invention also encompasses the above polynucleotide, further comprising a second nucleic acid. Preferably, the second nucleic acid encodes one or more polypeptides, and the physical location of the first nucleic acid relative to the second nucleic acid is such that, under the appropriate conditions, the first nucleic acid will operably regulate transcription of the second nucleic acid, thereby facilitating production of the one or more polypeptides. "Facilitation of production" includes increasing production constitutively, decreasing or blocking production except under specific conditions, and/or increasing production except under specific conditions. Among the "conditions" contemplated by the present invention are: (1) adding a component to a culture medium, (2) removing a component from a culture medium, (3) replacing one component of a culture medium with a second component, (4) increasing the temperature of the culture medium, (5) decreasing the temperature of the culture medium, and (6) regulating the atmospheric conditions (e.g., oxygen or nitrogen concentrations) in which the culture medium is maintained. Examples of such conditions are described in more detail, infra.

100851

One particularly preferred aspect of this embodiment is a polynucleotide comprising a nucleic acid, the sequence of which is at least 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98% or 99% identical to a nucleic acid sequence selected from the group consisting of SEQ ID NO:1, SEQ ID NO:2, SEQ ID NO:3, SEQ ID NO:4, SEQ ID NO:5, SEQ ID NO:6, SEQ ID NO:7, SEQ ID NO:8, SEQ ID NO:9, SEQ ID NO:10, SEQ ID NO:11, SEQ ID NO:12, SEQ ID NO:13, SEQ ID NO:14, SEQ ID NO:15, SEQ ID NO:16, SEQ ID NO:17, SEQ ID NO:18, SEQ ID NO:19, SEQ ID NO:20, SEQ ID NO:21, and SEQ ID NO:22; and a second nucleic acid, which encodes a component of an amino acid biosynthesis pathway. Examples of amino acid biosynthesis pathways included in this aspect of the invention are the enzymatic pathways that result in the synthesis of L-glycine, L-alanine, L-leucine, L-methionine, L-phenylalanine, L-tryptophan, L-lysine, L-glutamine, L-glutamic acid, L-aerginine, L-asparagine, L-aspartic acid, and L-threonine.

[0086]

A preferred amino acid biosynthesis pathway results in the synthesis of L-lysine. In this preferred aspect of this embodiment, the second nucleic acid encodes one of the following components of the L-lysine biosynthesis pathway: aspartokinase, aspartate beta-semialdehyde dehydrogenase, diaminopimelate dehydrogenase, diaminopimelate decarboxylase, dihydrodipicolinate synthetase, dihydrodipicolinate reductase, or pyruvate carboxylate.

[0087]

The present invention is further directed to a polynucleotide comprising a first nucleic acid, the sequence of which comprises about 6 contiguous nucleotides, preferably about 10 contiguous nucleotides, even more preferably about 12, 15, 20, 30, 40, 50, 100, 150, 200, or in some cases up to about 500, 1000, or 1500 contiguous nucleotides, of a sequence selected from the group consisting of SEQ ID NO:4, SEQ ID NO:5, SEQ ID NO:6, SEQ ID NO:7, SEQ ID NO:8, SEO ID NO:9, SEO ID NO:10, SEO ID NO:11, SEQ ID NO:12, SEQ ID NO:13, SEQ ID NO:14, SEQ ID NO:15, SEQ ID NO:16, SEQ ID NO:17, SEO ID NO:18, SEO ID NO:19, SEO ID NO:20, SEQ ID NO:21, and SEQ ID NO:22, or the complement of any of these sequences. By a first nucleic acid the sequence of which comprises about 10 contiguous nucleotides of any of said nucleic acid sequences, for example, is intended a nucleic acid which includes about 10 contiguous bases from any of said nucleotide sequences. Of course, the polynucleotide includes at least 10 contiguous nucleotides of any of said nucleic acid sequences, and may include about 12, 15, 20, 30, 40, 50, 100, 150, 200, or in some cases up to about 500, 1000, or 1500 contiguous nucleotides, or the entire sequence, of any of said nucleic acid sequences. In this context "about" includes the particularly recited size, larger or smaller by several (5, 4, 3, 2, or 1) nucleotides, at either terminus or at both termini.

[0088]

Representative examples of polynucleotides of the invention include a first nucleic acid the sequence of which comprises, for example, a sequence from about nucleotide 1-50, 51-100, 101-150, 151-200, 201-250, 251-300, 301-350, 351-400, 401-450, or 450-500 of SEQ ID NO:4, SEQ ID NO:5, SEQ ID NO:6, SEQ ID NO:7, SEQ ID NO:8, SEQ ID NO:10, SEQ ID NO:11, SEQ ID NO:13, SEQ ID NO:14, SEQ ID NO:15, SEQ ID NO:16, SEQ ID NO:17, SEQ ID NO:19, SEQ ID NO:20, SEQ ID NO:21, or SEQ ID NO:22, or the

complementary DNA strand thereto; a sequence from about nucleotide 1-50, 51-100, 101-150, 151-200, 201-250, 251-300, 301-350, 351-400, 401-450, 450-500, 501-550, 551-600, 601-650, 651-700, 701-750, 751-800, 801-850, 851-900, 901-850, 851-900, 901-850, 851-900, 901-850, 851-900, 901-850, 851-900, 901-850, 851-900, 901-850, 851-900, 901-850, 851-900, 901-850, 851-900, 901-850, 851-900, 901-850, 851-900, 901-850, 851-900, 901-850, 851-900, 901-850, 851-900, 901-850, 851-900, 901-850, 851-900, 901-850, 851-900, 901-850, 851-900, 901-850, 851-900, 901-850, 851-900, 901-850, 851-900, 901-850, 851-900, 901-850, 851-900, 901-850, 851-900, 901-850, 851-900, 901-850, 851-900, 901-850, 851-900, 901-850, 851-900, 901-850, 851-900, 901-850, 851-900, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 901-850, 9950, 951-1000, 1001-1050, 1051-1100, 1101-1150, 1151-1200, 1201-1250, 1251-1300, 1301-1350, 1351-1400, 1401-1450, 1451-1500 or 1501-1583 of SEO ID NO:9, or the complementary DNA strand thereto; a sequence from about $nucleotide\ 1-50, 51-100, 101-150, 151-200, 201-250, 251-300, 301-350, 351-400,\\$ 401-450, 450-500, 501-550, 551-600, 601-650, 651-700, 701-750, 751-800, 801-850, 851-900, 901-950, 951-1000, or 1001-1083 of SEQ ID NO:12, or the complementary DNA strand thereto; or a sequence from about nucleotide 1-50, 51-100, 101-150, 151-200, 201-250, or 251-297 of SEQ ID NO:18, or the complementary DNA strand thereto. In this context "about" includes the particularly recited ranges, larger or smaller by several (5, 4, 3, 2, or 1) nucleotides, at either terminus or at both termini. Additional examples of polynucleotides of the invention include a first nucleic acid the sequence of which comprises a consensus -10 or -35 region as listed in Table 2, infra.

[0089]

This embodiment of the present invention is directed to a polynucleotide comprising a first nucleic acid, the sequence of which comprises about 6 contiguous nucleotides, preferably about 10 contiguous nucleotides, even more preferably about 12, 15, 20, 30, 40, 50, 100, 150, 200, or in some cases up to about 500, 1000, or 1500 contiguous nucleotides, of a sequence selected from the group consisting of SEQ ID NO:4, SEQ ID NO:5, SEQ ID NO:6, SEQ ID NO:7, SEQ ID NO:8, SEQ ID NO:9, SEQ ID NO:10, SEQ ID NO:11, SEQ ID NO:12, SEQ ID NO:13, SEQ ID NO:14, SEQ ID NO:15, SEQ ID NO:16, SEQ ID NO:17, SEQ ID NO:18, SEQ ID NO:19, SEQ ID NO:20, SEQ ID NO:21, and SEQ ID NO:22, or the complement of any of these sequences, irrespective of whether they have functional activity. This is because even where a particular polynucleotide does not have functional activity, one of skill in the art would still know how to use the nucleic acid molecule, for instance, as a hybridization probe, an S1 nuclease mapping probe, or a polymerase chain reaction (PCR) primer.

[0090] Preferred, however, a polynucleotide comprising a first nucleic acid, the sequence of which comprises about 6 contiguous nucleotides, preferably about 10

contiguous nucleotides, even more preferably about 12, 15, 20, 30, 40, 50, 100, 150, 200, or in some cases up to about 500, 1000, or 1500 contiguous nucleotides, of a sequence selected from the group consisting of SEQ ID NO:4, SEQ ID NO:5, SEQ ID NO:6, SEQ ID NO:7, SEQ ID NO:8, SEQ ID NO:9, SEQ ID NO:10, SEQ ID NO:11, SEQ ID NO:12, SEQ ID NO:13, SEQ ID NO:14, SEQ ID NO:15, SEQ ID NO:16, SEQ ID NO:17, SEQ ID NO:18, SEQ ID NO:19, SEQ

ID NO:20, SEQ ID NO:21, and SEQ ID NO:22, or the complement of any of

these sequences, which do, in fact, have functional activity, in particular,

polynucleotides which regulate transcription in *Corynebacterium* species.

Examples of polynucleotides which regulate transcription in *Corynebacterium*

species include promoters, operators, transcription factor recognitions sites,

transcriptional complexes, and transcriptional regulatory elements, as described

herein.

known in the art.

[0091] Preferably, polynucleotides of the present invention comprise a promoter which functions in Corynebacterium species, preferably in Corynebacterium glutamicum, Brevibacterium flavum, or Brevibacterium lactofermentum, even more preferably in C. glutamicum. Assays to determine whether a polynucleotide is capable of regulating transcription in Corynebacterium species can be routinely performed using techniques described herein and otherwise

100921

Accordingly, the present invention also encompasses the above polynucleotide, further comprising a second nucleic acid. Preferably, the second nucleic acid encodes one or more polypeptides, and the physical location of the first nucleic acid relative to the second nucleic acid is such that, under the appropriate conditions, the first nucleic acid will operably regulate transcription of the second nucleic acid, thereby facilitating production of the one or more polypeptides, as described herein.

[0093]

One particularly preferred aspect of this embodiment is a polynucleotide comprising a first nucleic acid, the sequence of which comprises about 6 contiguous nucleotides, preferably about 10 contiguous nucleotides, even more preferably about 12, 15, 20, 30, 40, 50, 100, 150, 200, or in some cases up to about 500, 1000, or 1500 contiguous nucleotides, of a nucleic acid sequence selected from the group consisting of SEQ ID NO:1, SEQ ID NO:2, SEQ ID NO:3, SEQ ID NO:4, SEQ ID NO:5, SEQ ID NO:6, SEQ ID NO:7, SEQ ID NO:8, SEQ ID NO:9, SEQ ID NO:10, SEQ ID NO:11, SEQ ID NO:12, SEQ ID NO:13, SEQ ID NO:14, SEQ ID NO:15, SEQ ID NO:16, SEQ ID NO:17, SEQ ID NO:18, SEQ ID NO:19, SEQ ID NO:20, SEQ ID NO:21, and SEQ ID NO:22; and a second nucleic acid, which encodes a component of an amino acid biosynthesis pathway. A preferred amino acid biosynthesis pathway results in the synthesis of L-lysine. Examples of amino acid biosynthesis pathways, and specific components of the L-lysine biosynthesis pathway are described herein.

[0094]

The nucleotide sequences of functional bacterial promoters exhibit a limited but significant degree of variability depending on the gene that is regulated by the particular promoters in question, and the bacterial species in which the promoter is found. See, e.g., Pátek et al., Microbiology 142:1297 (1996), which is incorporated herein by reference in its entirety. Despite this variability, investigators have identified promoter consensus sequences by comparing the nucleotide sequences of multiple bacterial promoters to one another. A consensus sequence is a sequence that reflects the most common nucleotides that are found in particular positions for a multitude of promoters. Frequently, the functionality of a promoter consensus sequence is confirmed by mutational analysis in an experimental system; i.e., the expression of a reporter gene that is under the control of a mutated promoter is assessed.

[0095]

Early experiments and analyses of *E. coli* promoters identified a consensus sequence that comprises two nucleotide hexamers (six contiguous nucleotides) separated by about 17 nucleotides, and located about 35 and 10 nucleotides, respectively, upstream of the transcriptional start site for the

corresponding gene. Pátek et al., Microbiology 142:1297 (1996). These two hexamers are known individually as the -35 and -10 regions, and their respective consensus sequences are TTGaca, and TAtaaT. Within a consensus sequence, capital letters denote particular nucleotides that are found in analogous positions in at least 70% of the promoter sequences analyzed. Lower case letters denote nucleotides that are found in analogous positions in between 42% to 70% of the promoter sequences analyzed.

[0096]

Recently, Pátek et al. analyzed the sequences of 18 promoters from C. glutamicum, and four promoters from the temperate corynephage φ GA1, to identify a promoter consensus sequence specific to C. glutamicum. From their analysis, Pátek et al. identified a -35 consensus sequence of ttGcca, and a -10 consensus sequence of TA.aaT. The period (".") at the third position of the -10 consensus sequence indicates that the nucleotide at this position does not establish a consensus among the C. glutamicum promoters that were compared by Pátek et al.

[0097]

Accordingly, in one aspect of this embodiment of the present invention, a polynucleotide is provided which comprises a -10 promoter consensus sequence, and/or a -35 consensus sequence derived from a sequence selected from the group consisting of SEQ ID NO:4, SEQ ID NO:5, SEQ ID NO:6, SEQ ID NO:7, SEQ ID NO:8, SEQ ID NO:9, SEQ ID NO:10, SEQ ID NO:11, SEQ ID NO:12, SEQ ID NO:13, SEQ ID NO:14, SEQ ID NO:15, SEQ ID NO:16, SEQ ID NO:17, SEQ ID NO:18, SEQ ID NO:19, SEQ ID NO:20, SEQ ID NO:21, and SEQ ID NO:22. While not being bound by theory, such sequence motifs may function as part of a promoter capable of regulating transcription in Corynebacterium species, preferably in Corynebacterium glutamicum, Brevibacteriumflavum, or Brevibacterium lactofermentum, even more preferably in C. glutamicum.

[0098]

To identify -10 promoter and/or -35 promoter consensus sequences, computer analyses of SEQ ID NOs: 1 through 22 were conducted using the prokaryotic nucleic acid motif search functionality provided by the OMIGA 2.0 computer software package. In addition, SEQ ID NOs: 1 through 22 were analyzed visually to identify any sequences that agree with the consensus sequences identified by Pátek et al. A summary of predicted -10 and -35 consensus sequences identified for SEQ ID NO: 1 through 22 is listed in Table 2. It is to be noted that in the numbering convention of SEQ ID NOs 1 through 22, all nucleotides have a positive number and the numbers increase. This is in contrast to the normal convention for promoter regions, wherein the nucleotides are assigned negative numbers which increase up to nucleotide number 1 at the putative transcription start site.

TABLE 2

SEQ ID NO:	-10 sequence(s)	Position	-35 sequence(s)	Position
1			TTGAGA*	375
1			TTGAAA*	386
3			TTGAAA*	339
4	TACAAT [‡]	275		
4			TTGAGA*	57
4			TTGAAA*	186
4			TTGAAA*	301
4			TTGCCA [‡]	466
5			TTGAAA*	307
5			TTGAGA*	340
7	TACAAT‡	334		
7			TTGCCA‡	224
8			TTGATA*	341
8			TTGCCA [‡]	371
9			TTGATA*	803
10	TACAAT [‡]	406		
10			TTGACA*	234
11			TTGAAA*	248
13	TACAAT [‡]	394		
13			TTGCCA [‡]	118
14	TAAAAT‡	169		
15	TAAAAT‡	379		
17			TTGACA*	211
18			TTGAAA*	31
18			TTGAAA*	65
18	TAAAAT [‡]	269		
18			TTGCCA [‡]	168
20			TTGAAA*	490
21			TTGAAA*	41
21			TTGAGA*	445
21			TTGAAA*	494

^{*} consensus sequences identified by the OMIGA 2.0 program

 $[\]updownarrow$ consensus sequences identified visually by comparison to consensus sequences set forth in Pátek et al.

[0099]

In another embodiment, the invention provides an isolated polynucleotide comprising a first nucleic acid at least 10 nucleotides in length, preferably at least 12, 15, 30, 50 or 150 nucleotides in length, which hybridizes under stringent conditions to a reference nucleic acid, or the complement thereof, wherein the sequence of said reference nucleic acid is selected from the group consisting of SEQ ID NO:4, SEQ ID NO:5, SEQ ID NO:6, SEQ ID NO:7, SEQ ID NO:8, SEQ ID NO:9, SEQ ID NO:10, SEQ ID NO:11, SEQ ID NO:12, SEQ ID NO:13, SEQ ID NO:14, SEQ ID NO:15, SEQ ID NO:16, SEQ ID NO:17, SEQ ID NO:18, SEQ ID NO:19, SEQ ID NO:20, SEQ ID NO:21, and SEQ ID NO:22. By "stringent hybridization conditions" is intended overnight incubation at 42°C in a solution comprising: 50% formamide, 5x SSC (750 mM NaCl, 75mM trisodium citrate), 50 mM sodium phosphate (pH 7.6), 5x Denhardt's solution, 10% dextran sulfate, and 20 μg/ml denatured, sheared salmon sperm DNA, followed by washing the filters in 0.1x SSC at about 65°C.

[0100]

By a first nucleic acid "at least 10 nucleotides in length," which hybridizes to a reference nucleic acid is intended a first nucleic acid (either DNA or RNA) which hybridizes to 10 or more contiguous nucleotides from the nucleotide sequence of the reference nucleic acid (e.g., SEQ ID NOs 4-22).

[0101]

This embodiment of the present invention is directed to an isolated polynucleotide comprising a first nucleic acid at least 10 nucleotides in length, preferably at least 12, 15, 30, 50 or 150 nucleotides in length, which hybridizes under stringent conditions to a reference nucleic acid, or the complement thereof, wherein the sequence of said reference nucleic acid is selected from the group consisting of SEQ ID NO:4, SEQ ID NO:5, SEQ ID NO:6, SEQ ID NO:7, SEQ ID NO:8, SEQ ID NO:9, SEQ ID NO:10, SEQ ID NO:11, SEQ ID NO:12, SEQ ID NO:13, SEQ ID NO:14, SEQ ID NO:15, SEQ ID NO:16, SEQ ID NO:17, SEQ ID NO:18, SEQ ID NO:19, SEQ ID NO:20, SEQ ID NO:21, and SEQ ID NO:22, or the complement of any of these sequences, irrespective of whether they have functional activity. This is because even where a particular polynucleotide does not have functional activity, one of skill in the art would still know how to

[0102]

use the nucleic acid molecule, for instance, as a hybridization probe, an S1 nuclease mapping probe, or a polymerase chain reaction (PCR) primer.

Preferred, however, an isolated polynucleotide comprising a first nucleic acid at least 10 nucleotides in length, preferably at least 12, 15, 30, 50 or 150 nucleotides in length, which hybridizes under stringent conditions to a reference nucleic acid, or the complement thereof, wherein the sequence of said reference nucleic acid is selected from the group consisting of SEQ ID NO:4, SEQ ID NO:5, SEQ ID NO:6, SEQ ID NO:7, SEQ ID NO:8, SEQ ID NO:9, SEQ ID NO:10, SEQ ID NO:11, SEQ ID NO:12, SEQ ID NO:13, SEQ ID NO:14, SEQ ID NO:15, SEQ ID NO:16, SEQ ID NO:17, SEQ ID NO:18, SEQ ID NO:19, SEQ ID NO:20, SEQ ID NO:21, and SEQ ID NO:22, or the complement of any of these sequences, which do, in fact, have functional activity, in particular, polynucleotides which regulate transcription in Corynebacterium species. Examples of polynucleotides which regulate transcription in Corynebacterium species include promoters, operators, transcription factor recognitions sites, transcriptional complexes, and transcriptional regulatory elements, as described herein. Preferably, polynucleotides of the present invention comprise a promoter which functions in Corynebacterium species, preferably in Corynebacterium glutamicum, Brevibacterium flavum, or Brevibacterium lactofermentum, even Assays to determine whether a more preferably in C. glutamicum. polynucleotide is capable of regulating transcription in Corynebacterium species can be routinely performed using techniques described herein and otherwise known in the art.

[0103] Accordingly, the present invention also encompasses the above polynucleotide, further comprising a second nucleic acid. Preferably, the second nucleic acid encodes one or more polypeptides, and the physical location of the first nucleic acid relative to the second nucleic acid is such that, under the appropriate conditions, the first nucleic acid will operably regulate transcription of the second nucleic acid, thereby facilitating production of the one or more polypeptides, as described herein.

[0104] One particularly preferred aspect of this embodiment is an isolated polynucleotide comprising a first nucleic acid at least 10 nucleotides in length, preferably at least 12, 15, 30, 50 or 150 nucleotides in length, which hybridizes under stringent conditions to a reference nucleic acid, or the complement thereof, wherein the sequence of said reference nucleic acid is selected from the group consisting of selected from the group consisting of SEQ ID NO:1, SEQ ID NO:2, SEQ ID NO:3, SEQ ID NO:4, SEQ ID NO:5, SEQ ID NO:6, SEQ ID NO:7, SEQ ID NO:8, SEQ ID NO:9, SEQ ID NO:10, SEQ ID NO:11, SEQ ID NO:12, SEQ ID NO:13, SEQ ID NO:14, SEQ ID NO:15, SEQ ID NO:16, SEQ ID NO:17, SEQ ID NO:18, SEQ ID NO:19, SEQ ID NO:10, SEQ ID NO:21, and SEQ ID NO:22; and a second nucleic acid, which encodes a component of an amino acid biosynthesis pathway. A preferred amino acid biosynthesis pathway results in the synthesis of L-lysine. Examples of amino acid biosynthesis pathways, and

[0105]

Another embodiment of the invention provides an isolated polynucleotide comprising a first nucleic acid, the sequence of which is at least 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, or 99% identical to SEQ ID NO: 2, and a second nucleic acid which encodes a polypeptide which functions as a component of the lysine biosynthesis pathway. Alternatively in this embodiment, the sequence of the first nucleic acid comprises about 6 contiguous nucleotides, preferably about 10 contiguous nucleotides, even more preferably about 12, 15, 20, 30, 40, 50, 100, 150, 200, or 500, contiguous nucleotides, of SEQ ID NO:2, or the first nucleic acid is at least 10 nucleotides in length, preferably at least 12, 15, 30, 50 or 150 nucleotides in length, and hybridizes under stringent conditions to a reference nucleic acid, or the complement thereof, wherein the sequence of said reference nucleic acid is SEQ ID NO:2. A preferred amino acid biosynthesis pathway results in the synthesis of L-lysine. Examples of components of the Llysine biosynthesis pathway are disclosed herein. In this embodiment, the first nucleic acid, which comprises the C. glutamicum aceA promoter, regulates transcription of the second nucleic acid.

specific components of the L-lysine biosynthesis pathway are disclosed herein.

[0106]

Yet another embodiment of the invention provides an isolated polynucleotide comprising a first nucleic acid, the sequence of which is at least 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, or 99% identical to SEQ ID NO: 3; and a second nucleic acid which encodes polypeptide which functions as a component of an amino acid biosynthesis pathway. Alternatively in this embodiment, the sequence of the first nucleic acid comprises about 6 contiguous nucleotides, preferably about10 contiguous nucleotides, even more preferably about 12, 15, 20, 30, 40, 50, 100, 150, 200, or 500, contiguous nucleotides, of SEQ ID NO:3, or the first nucleic acid is at least 10 nucleotides in length, preferably at least 12, 15, 30, 50 or 150 nucleotides in length, and hybridizes under stringent conditions to a reference nucleic acid, or the complement thereof, wherein the sequence of said reference nucleic acid is SEQ ID NO:3. A preferred amino acid biosynthesis pathway results in the synthesis of L-lysine. Examples of amino acid biosynthesis pathways, and specific components of the L-lysine biosynthesis pathway are disclosed herein. In this embodiment, the first nucleic acid, which comprises the C. glutamicum aceB promoter, regulates transcription of the second nucleic acid.

[0107]

A further embodiment of the invention provides an isolated Corynebacterium chromosome with a first nucleic acid integrated into the chromosome, the sequence of which is at least 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, or 99% identical to SEQ ID NO:1. By "an isolated Corynebacterium species chromosome" is meant a bacterial chromosome which has been altered in some way to affect gene expression, either of naturally-occuring genes or heterologous genes, and then isolated from other non-altered chromosome. In this embodiment, the isolated Corynebacterium species chromosome also has a second nucleic acid integrated therein, wherein said second nucleic acid encodes polypeptide which functions as a component of an amino acid biosynthesis pathway results in the synthesis of L-lysine. Examples of amino acid biosynthesis

pathways, and specific components of the L-lysine biosynthesis pathway are disclosed herein. In this embodiment, the first nucleic acid, which comprises the C. glutamicum pta promoter, regulates transcription of the second nucleic acid.

[0108] Another embodiment of the invention provides an isolated Corynebacterium chromosome with a first nucleic acid integrated into the chromosome, the sequence of which is at least 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, or 99% identical to SEQ ID NO:23. In this embodiment, the isolated Corynebacterium species chromosome also has a second nucleic acid integrated therein, wherein said second nucleic acid encodes polypeptide which functions as a component of an amino acid biosynthesis pathway, preferably an L-lysine biosynthesis pathway. In this embodiment, the first nucleic acid, which comprises the E. coli araBAD promoter and the araC regulator molecule, regulates transcription of the second nucleic acid. The araC/araBAD transcriptional complex is a tightly regulated system responsive to the regulator molecule arabinose.

Yet another embodiment of the invention provides an isolated [0109]Corynebacterium chromosome with a first nucleic acid integrated into the chromosome, the sequence of which is identical to either SEQ ID NO:26 or SEQ As with the embodiments described above, the isolated ID NO:27. Corynebacterium species chromosome also has a second nucleic acid integrated therein, wherein said second nucleic acid encodes polypeptide which functions as a component of an amino acid biosynthesis pathway, preferably an L-lysine biosynthesis pathway, and the first nucleic acid, which is a trc or tac synthetic promoter, regulates transcription of the second nucleic acid. These synthetic promoters combine the -35 region from the trp promoter with the -10 region from the lacUV5 promoter. See Brosius, J. et al., J. Biol. Chem. 260, 3539 (1985). In the embodiment, the isolated chromosome further has a third nucleic acid integrated therein, the sequence of which is at least 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, or 99% identical to SEQ ID NO:28. This third nucleic acid is operably linked to a transcription control region, and the nucleic acid encodes a polypeptide at least 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, or 99% identical to SEQ ID NO:29, the *E. coli lac* repressor gene. The lacf^a-trc (SEQ ID NO:30) or the lacf^a-trac transcriptional complex is responsive to the regulatory molecules isoproplythiogalactoside (IPTG) and lactose. This regulatory complex in *Corynebacterium glutamicum*, however, is leaky, *i.e.*, transcription occurs in the absence of any added inducer.

[0110] In each of the isolated chromosomes described above, the second nucleic acid may be a naturally occurring Corynebacterium nucleic acid, situated in its normal position on the chromosome, it may be a naturally occurring Corynebacterium nucleic acid which has been moved to a different position on the chromosome, or it may be a non-native nucleic acid. The first nucleic acid is engineered to be in a physical location which allows it to regulate transcription of the second nucleic acid either through in vitro cloning of a vector having both the first and second nucleic acids followed by homologous or random recombination into the Corynebacterium species chromosome, or it is inserted in front of a selected second nucleic acid already situated in the chromosome through homologous recombination. Such manipulations are readily understood by those of ordinary skill in the art.

[0111] In another aspect of the above embodiments, a Corynebacterium species host cell comprising a Corynebacterium species chromosome as described above is provided, preferably a host cell derived from Corynebacterium glutamicum, Brevibacterium flavum, and Brevibacterium lactofermentum, and even more preferably derived from Corynebacterium glutamicum. In a related aspect, a method of producing such a Corynebacterium species host cell is provided, which comprises transforming Corynebacterium species cells with a vector comprising a first nucleic acid as described above, wherein said vector facilitates integration of the first nucleic acid into the chromosome of said Corynebacterium species cells, and selecting the host cell in certain preferred embodiments, the vector also comprises the second nucleic acid described above, and/or the third nucleic acid

described above, with the first nucleic acid being physically situated to regulate transcription of the second nucleic acid. These vectors are also provided.

[0112] The present invention further relates to variants of the polynucleotides of the present invention. Variants may occur naturally, such as a natural allelic variant. By an "allelic variant" is intended one of several alternate forms of a gene occupying a given locus on a chromosome of an organism. *Genes II*, Lewin, B., ed., John Wiley & Sons, New York (1985). Non-naturally occurring variants may be produced using art-known mutagenesis techniques.

[0113] Such variants include those produced by nucleotide substitutions, deletions or additions which may involve one or more nucleotides. Particularly preferred are variants which alter the level of activity of the polynucleotide relative to a non-variant polynucleotide. For example, variants which increase or decrease the strength of a promoter, or variants which alter or eliminate the need for an inducer. Methods to construct variant promoters and screen them for their relative strengths utilizing reporter genes are well known in the art. See, e.g., Vašicová, et al., J. Bacteriol. 181:6188-6199 (1999), which is incorporated herein by reference in its entirety.

D Vectors and Host Cells

- [0114] The present invention also relates to vectors which include polynucleotides of the present invention, host cells which are engineered with vectors of the invention and methods to increase the production of metabolic products such as amino acids purines, pyrimidines, and heterologous polypeptides through utilization of host cells of the invention.
- [0115] Host cells can be engineered to incorporate polynucleotides of the present invention. The polynucleotides may be introduced alone or with other polynucleotides. Such other polynucleotides may be introduced independently, co-introduced or introduced joined to the polynucleotides of the invention.

[0116] In accordance with this aspect of the invention the vector may be, for example, a plasmid vector or a bacteriophage vector. Such vectors may be introduced into Corynebacterium cells such that they are maintained and replication as an extrachromosomal element, or, alternatively, such that they integrate into the chromosome. Vectors are introduced into host cells by well known techniques for introducing DNA and RNA into prokaryotic cells, examples of which are described in the Examples, infra.

[0117] Preferred vectors comprise a first nucleic acid which is capable of regulating transcription of a second nucleic acid which is inserted in operable association, preferably immediately downstream, of the first nucleic acid. The second nucleic acid preferably encodes a polypeptide such as an enzyme in an amino acid biosynthesis pathway. Especially preferred among vectors are those which allow convenient insertion of the second nucleic acid into the vector into the proper position to be regulated by the first nucleic acid, for example, through the use of a multiple cloning region. Appropriate trans-acting factors such as inducers either are supplied by the host cell, supplied by a complementing vector or supplied by the vector itself upon introduction into the host cell.

[0118] A great variety of vectors can be used in the invention. Such vectors include chromosomal, episomal and virus-derived vectors e.g., vectors derived from bacterial plasmids and from bacteriophage, as well as vectors derived from combinations thereof, such as those derived from plasmid and bacteriophage genetic elements, such as cosmids and phagemids, all may be used in accordance with this aspect of the present invention. Generally, any vector suitable to maintain and propagate a polynucleotide in a bacterial host may be used in this regard.

[0119] A large numbers of suitable vectors for use in bacteria are known, many of which are commercially available. Preferred prokaryotic vectors include plasmids such as those capable of replication in E. coli (such as, for example, pBR322, ColEl, pSC101, pACYC 184, πVX. Such plasmids are, for example, disclosed by Maniatis, T., et al., In: Molecular Cloning, A Laboratory Manual,

Cold Spring Harbor Press, Cold Spring Harbor, NY (1982)). The following vectors are provided by way of example: pET (Novagen), pQE70, pQE60, pQE-9 (Qiagen), pBs, phagescript, psiXI74, pBlueScript SK, pBsKS, pNH8a, pNH16a, pNH18a, pNH46a (Stratagene); pTrc99A, pKK223-3, pKK233-3, pDR540, pRIT5 (Pharmacia).

- [0120] Preferred vectors for the isolated polynucleotides of the invention include the pFC1 to pFC7 novel family of combinatorial cloning vectors (Lonsdale, D.M., et al., Plant Molecular Biology Reporter 13: 343-345 (1995)), the pK184 vector (Jobling, M.G. and Homes, R.K., Nucleic Acid Research 18: 5315-5316 (1990)).
- [0121] Another group of preferred vectors are those that are capable of autonomous replication in *Corynebacterium* species. Such vectors are well known to those skilled in the art of amino acid production by way of microbial fermentation, examples of which include pSR1, pMF1014α and vectors derived therefrom.
- [0122] The second nucleic acid encoding a polypeptide is operatively associated with the first nucleic acid as described above, which provides the appropriate transcription control sequence(s)), in particular, a promoter. In addition, the vector may contain sites for transcription initiation and termination, and, in the transcribed region, a ribosome binding site for translation. The second nucleic acid, which provides the coding region of the polypeptide to be regulated will include a translation initiating AUG or GUG at the beginning and a termination codon (UAA, UGA or UAG) appropriately positioned at the end of the coding region.
- [0123] In addition, and as described in more detail herein, the vector may contain control regions that regulate as well as engender expression. Generally, such regions will operate by controlling transcription, such as inducer or repressor binding sites and enhancers, among others.
- [0124] Vectors of the present invention generally will include a selectable marker. Such markers also may be suitable for amplification or the vectors may contain additional markers for this purpose. In this regard, vectors preferably

contain one or more selectable marker genes to provide a phenotypic trait for selection of transformed host cells. Such markers include, but are not limited to, an antibiotic resistance gene such as a chloramphenicol, ampicillin, or kanamycin resistance gene, or an autotrophic gene which allows the host cell to grow in the absence of a nutrient for which the host cell strain is normally autotrophic.

- [0125] If the vector is intended to be maintained in the host cell extrachromosomally, it will contain, in addition and origin of replication which will allow it to replicate in the Corynebacterium species host cell. Alternatively, if it is desired that the vector integrate into the Corynebacterium species chromosome, the vector is constructed such that it cannot replicate in Corynebacterium. For example, such a vector might be capable of propagation in another organism, for example, E. coli, but lack the proper origin of replication to be propagated in Corynebacterium. In another aspect of this embodiment, the vector is a shuttle vector which can replicate and be maintained in more than one host cell species, for example, such a shuttle vector might be capable of replication in a Corynebacterium host cell such as a C. glutamicum host cell, and also in an E. coli host cell.
- [0126] A vector comprising a polynucleotide as described elsewhere herein may be introduced into an appropriate Corynebacterium species host cell using a variety of well known techniques, for example, transformation or electroporation. Appropriate culture mediums and conditions for culturing Corynebacterium host cells are known in the art.
- [0127] The invention provides methods for increasing amino acid production and processes for the production of an amino acid wherein increased biosynthetic pathway gene expression is accomplished through insertion of an isolated polynucleotide into the chromosome of a host cell. For example, insertion of isolated polynucleotides into the chromosome of Corynebacterium species may be done utilizing the pK184 plasmid described by Jobling, M. et al., Nucleic Acids Research 18(17): 5315-5316 (1990). Because these vectors lack a Corynebacterium species origin of replication and contains a selectable marker

such as kanamycin (kan), cells will only be capable of growing under selection if the vector has been inserted into the host cell chromosome by homologous recombination.

- [0128] The invention also provides methods for increasing amino acid production and processes for the production of an amino acid wherein increased biosynthetic pathway gene expression is accomplished through the introduction into a host cell of a self-replicating, extra-chromosomal vector, e.g., a plasmid, comprising an isolated nucleic acid molecule encoding an amino acid biosynthetic pathway gene or genes. Suitable plasmids for these embodiments include pSR1 and other derivatives of pSR1 (Archer, J. et al., J. Gen. Microbiol. 139: 1753-1759 (1993)).
- [0129] These vectors are listed solely by way of illustration of the many vectors available to those of skill in the art. Other vectors are described in the Examples, infra. Selection of appropriate vectors to transform into a host cell is a well known procedure and the requisite techniques for vector construction, introduction of the vector into the host and maintenance in the host are routine skills in the art.
- [0130] The present invention also relates to Corynebacterium species host cells comprising the above-described vector constructs described herein. The host cell can be any Corynebacterium species, preferably Corynebacterium glutamicum, Brevibacteriumflavum, or Brevibacteriumlactofermentum, even more preferably a strain of C. glutamicum. Preferred strains of C. glutamicum to use as host cells of the invention include: NRRL-B11474, ATCC 21799, ATCC 21529, ATCC 21543, and E12.
- [0131] Introduction of a vector into the host cell can be effected by transformation, electroporation, transduction, infection or other methods. Such methods are described in many standard laboratory manuals, such as Davis et al., Basic Methods in Molecular Biology (1986).

F. Methods and Processes of the Invention

[0132] Various embodiments of the invention provide methods to increase the production of an amino acid and processes for the production of an amino acid from a Corynebacterium species host cell. Particularly preferred Corynebacterium species of the methods and processes of the invention include: Corynebacterium glutamicum, Brevibacterium flavum, Brevibacterium lactofermentum and other Cornynebacteria and Brevibacteria species known in the art.

[0133] Amino acid biosynthetic pathway genes embodied by the methods and processes described herein include those for L-glycine, L-alanine, L-methionine, L-phenylalanine, L-tryptophan, L-proline, L-serine, L-threonine, L-cysteine, L-tyrosine, L-asparagine, L-glutamine, L-aspartic acid, L-glutamic acid, L-lysine, L-arginine, L-histidine, L-isoleucine, L-leucine, and L-valine biosynthesis. Particularly preferred embodiments are drawn to biosynthetic pathway genes for L-lysine (Sahm et al., Ann. N. Y. Acad. Sci. 782: 25-39 (1996)), L-threonine, L-isoleucine, L-tryptophan, and L-valine.

[0134] By way of example, the amino acid pathway for L-lysine biosynthesis is well known to skilled artisans of amino acid production in Corynebacterium species. Genes encoding the enzymes important for the conversion of L-aspartate to L-lysine include the ask, asd, dapA, dapB, ddh, and lysA genes, which encode, respectively, aspartokinase, aspartate beta-semialdehyde dehydrogenase, dihydrodipicolinate synthetase, dihydrodipicolinate reductase, diaminopimelate dehydrogenase, diaminopimelate decarboxylase, and pyruvate carboxylase. Thus, the invention provides herein for exemplary purposes only, specific embodiments utilizing L-lysine biosynthetic pathway genes. Other embodiments drawn to the use of biosynthetic pathway genes for the synthesis of other amino acids are also encompassed by the invention described herein.

[0135] The methods to increase the production of a biosynthetic product such as an amino acid and the processes for the production of the biosynthetic product of

the invention involve increasing the expression of at least one biosynthetic pathway gene. Preferred methods of increasing expression comprise using promoters of the invention.

Accordingly, the present invention provides a first nucleic acid which [0136] preferably comprises a promoter. Preferred promoters include, but are not limited to the following types of promoters: a constitutive promoter, an inducible promoter, a derepressable promoter, a heat sensitive promoter, and a cold sensitive promoter. In this aspect of the invention, a constitutive promoter causes the second nucleic acid to be transcribed at a substantially constant rate; an inducible promoter will be non-functional or will function at a reduced level under certain standard culture conditions, but causes elevated transcription of the second nucleic acid when the transformed host cells are grown in the presence of an appropriate inducer substance; a derepressable promoter causes elevated transcription of the second nucleic acid when the transformed host cells are grown in the absence of an appropriate suppressor substance, but will be nonfunctional or will function at a reduced level when the suppressor substance is present; a heat sensitive promoter causes elevated transcription of the second nucleic acid when the transformed host cells are grown in medium the temperature of which is adjusted to a temperature that is greater than the optimum temperature for wild-type untransformed host cells; and a cold sensitive promoter causes elevated transcription of the second nucleic acid when said transformed host cells are grown in a medium the temperature of which is adjusted to a temperature that is less than the optimum temperature for a wild-type untransformed host cell.

[0137] Also included within this embodiment of the invention are methods wherein the transformed host cells are grown to a sufficient cell density according to methods well known in the art. Further included in this embodiment are methods for substantially purifying the biosynthetic product from said culture, wherein said purification methods are also well known in the art. [0138] In certain embodiments of the present invention, an inducible promoter is used. When an inducible promoter is used, it is preferred that the transformed Corynebacterium species host cells are first allowed to proliferate to a sufficient cell density in the absence of the inducer, followed by addition of the inducer to the culture medium thereby stimulating production of said product. Examples of inducers include, but are not limited to one or more of the following substances: acetic acid, pyruvate, ethanol, fatty acids, cellulose subunits, starch subunits, triglycerides, or any of the following sugars: fructose, maltose, lactose, or ambinose.

[0139] In other embodiments, a derepressable promoter is used. When a derepressable promoter is used, it is preferred that the transformed *Corynebacterium* species host cells are first allowed to proliferate to a sufficient cell density in the presence of the suppressor, followed by either removal of the suppressor from the culture, or replacement of the medium containing the suppressor with medium lacking the suppressor, thereby stimulating production of said product. Under certain conditions, the suppressor is removed from the medium by being metabolized by the host cell. Examples of said suppressors include, but are not limited to any of the following substances: a purine, a pyrimidine, oxygen, or a suppressor amino acid including histidine.

[0140] In other embodiments, the promoter is self-inducing, i.e., the inducer molecule is produced by the host cell and builds up through the course of the growth, thereby changing gene expression as growth progresses. In this embodiment, the promoter is induced either gradually or at some point in the growth of the host cell culture, however this induction does not require the addition of exogenous compounds to the culture medium.

[0141] In yet another embodiment, a heat sensitive promoter is used. When a heat sensitive promoter is used, it is preferred that the transformed host cells are first allowed to proliferate to a sufficient cell density in culture medium, the temperature of which is adjusted to the optimum growth temperature for untransformed host cells, followed by adjusting the temperature of said culture

medium to a temperature that is greater than the optimum growth temperature for untransformed host cells.

[0142] In yet another embodiment, a cold sensitive promoter is used. When a cold sensitive promoter is used, it is preferred that the transformed host cells are first allowed to proliferate to a sufficient cell density in culture medium, the temperature of which is adjusted to the optimum growth temperature for untransformed host cells, followed by adjusting the temperature of said culture medium to a temperature that is less than the optimum growth temperature for untransformed host cells.

In a related embodiment of the invention, a method is provided for the production of a product such as an amino acid, a purine nucleotide, or a heterologous polypeptide in a *Corynebacterium* species host cell utilizing a heterologous transcriptional complex. According to this embodiment, the host cell harbors the heterologous transcriptional complex integrated into the bacterial chromosome. Preferred heterologous transcriptional complexes include the *E. coli ara*C-*ara*BAD transcriptional complex (SEQ ID NO:23), and the lacf^a-trc or lacf^a-tac transcriptional complexes which combine the *E. coli lac*f^a repressor gene (lacf^a, SEQ ID NO:28) with either the trc promoter (SEQ ID NO:26 or SEQ ID NO:30) or the tac promoter (SEQ ID NO:27). These synthetic promoters combine the -35 region from the trp promoter with the -10 region from the lacUV5 promoter. See Brosius, J. et al., J. Biol. Chem. 260, 3539 (1985).

[0144] The method utilizing the E. coli araC-araBAD transcriptional complex preferably further comprises culturing the host cell in or on a medium to which the inducer arabinose has been added, either throughout growth of the host cells, or at a point during growth where the host cell culture has reached an optimal density. The method utilizing either the lacl^q-trc or the lacl^q-tac transcriptional complex may further comprise culturing the host cell in or on a medium to which the inducer isoproplythiogalactoside (IPTG) or the inducer lactose has been added, either throughout growth of the host cells, or at a point during growth where the host cell culture has reached an optimal density. However, in C.

glutamicum, the lacI^q/trc system is leaky, therefore an inducer is not required for optimal growth.

[0145] A preferred aspect of this embodiment provides a method for the production of an amino acid in a Corynebacterium host cell utilizing a heterologous transcriptional complex integrated into the host cell chromosome, where the heterologous transcriptional complex operably regulates production a polypeptide which functions in an amino acid biosynthesis pathway. In this aspect, the heterologous transcriptional complex, when integrated into the host cell chromosome, regulates transcription of one or more metabolic pathway enzymes, thereby resulting in elevated production of the amino acid by the transformed host cell.

[0146] One aspect of this embodiment involves addition of an inducer to the culture medium thereby upregulating the heterologous transcriptional complex. In the method utilizing the *E. coli araC-araBAD* transcriptional complex arabinose is added to the medium. In the method utilizing the lacl^q-trc or the lacl^q-trc transcriptional complex, IPTG and/or lactose is added to the culture medium. Preferably in this method, the host cells are grown to a sufficient cell density according to methods well known in the art prior to addition of the inducer molecule.

[0147] For those embodiments of the invention drawn to a method to increase production of an amino acid, screening for increased production of an amino acid, for example L-lysine, may be determined by directly comparing the amount of L-lysine produced in culture by a standard Corynebacterium species strain to that of a Corynebacterium species host cell which has been transformed to comprise an isolated polynucleotide of the present invention. The level of production of the amino acid of choice may conveniently determined by the following formula to calculate the percent yield from dextrose: [(g amino acid/L / (g dextrose consumed/L)]*100.

[0148] A variety of media known to those skilled in the art may be used to support cell growth for the production of an amino acid. Illustrative examples of suitable carbon sources include, but are not limited to: carbohydrates, such as glucose, fructose, sucrose, starch hydrolysate, cellulose hydrolysate and molasses; organic acids, such as acetic acid, propionic acid, formic acid, malic acid, citric acid, and fumaric acid; and alcohols, such as glycerol. Illustrative examples of suitable nitrogen sources include, but are not limited to: ammonia, including ammonia gas and aqueous ammonia; ammonium salts of inorganic or organic acids, such as ammonium chloride, ammonium phosphate, ammonium sulfate and ammonium acetate; and other nitrogen-containing sources, including meat extract, peptone, corn steep liquor, casein hydrolysate, soybean cake hydrolysate, urea and yeast extract.

[0149]

A variety of fermentation techniques are known in the art which may be employed in processes of the invention drawn to the production of amino acids. Generally, amino acids may be commercially produced from the invention in fermentation processes such as the batch type or of the fed-batch type. In batch type fermentations, all nutrients are added at the beginning of the fermentation. In fed-batch or extended fed-batch type fermentations one or a number of nutrients are continuously supplied to the culture, right from the beginning of the fermentation or after the culture has reached a certain age, or when the nutrient(s) which are fed were exhausted from the culture fluid. A variant of the extended batch of fed-batch type fermentation is the repeated fed-batch or fill-and-draw fermentation, where part of the contents of the fermenter is removed at some time, for instance when the fermenter is full, while feeding of a nutrient is continued. In this way a fermentation can be extended for a longer time.

[0150]

Another type of fermentation, the continuous fermentation or chemostat culture, uses continuous feeding of a complete medium, while culture fluid is continuously or semi-continuously withdrawn in such a way that the volume of the broth in the fermenter remains approximately constant. A continuous fermentation can in principle be maintained for an infinite time.

[0151]

In a batch fermentation an organism grows until one of the essential nutrients in the medium becomes exhausted, or until fermentation conditions become unfavorable (e.g. the pH decreases to a value inhibitory for microbial growth). In fed-batch fermentations measures are normally taken to maintain favorable growth conditions, e.g. by using pH control, and exhaustion of one or more essential nutrients is prevented by feeding these nutrient(s) to the culture. The microorganism will continue to grow, at a growth rate dictated by the rate of nutrient feed. Generally a single nutrient, very often the carbon source, will become limiting for growth. The same principle applies for a continuous fermentation, usually one nutrient in the medium feed is limiting, all other nutrients are in excess. The limiting nutrient will be present in the culture fluid at a very low concentration, often unmeasurably low. Different types of nutrient limitation can be employed. Carbon source limitation is most often used. Other examples are limitation by the nitrogen source, limitation by oxygen, limitation by a specific nutrient such as a vitamin or an amino acid (in case the microorganism is auxotrophic for such a compound), limitation by sulphur and limitation by phosphorous.

[0152] The amino acid may be recovered by any method known in the art. Exemplary procedures are provided in the following: Van Walsem, H.J. & Thompson, M.C., J. Biotechnol. 59:127-132 (1997), and U.S. Pat. No. 3,565,951, both of which are incorporated herein by reference.

EXAMPLES

EXAMPLE 1

Production of Host Cells Utilizing Transcriptional Regulatory Regions of the

Present Invention

[0153] Corynebacterium species host cells, in which to express metabolic polypeptides are constructed as follows. Polynucleotides of the present invention which comprise transcriptional regulatory regions related to SEQ ID NOs 1-23, 26-28, and 30 are inserted upstream of a multiple cloning region, thereby allowing convenient insertion coding regions to be regulated and expressed. Testing of the various transcriptional regulatory regions is conveniently carried by inserting the various regulatory regions in operable association to a known reporter gene, such as β-galactosidase (*lacZ*). Methods and techniques common to the art of recombinant DNA technology are used in making vectors of the invention, as may be found in the many laboratory manuals cited and incorporated herein, for example as found in J. Sambrook, E.F. Fritsch and T. Maniatis, *Molecular Cloning: A Laboratory Manual*, 2d ed., Cold Spring Harbor Laboratory Press, Cold Spring Harbor, New York (1989). Construction of exemplary vectors is set forth in Example 2, *infra*.

[0154]

For cell transformation experiments with the isolated nucleic acid molecules of the invention, the growth and preparation of competent cells may be done according to the following procedure: (1) picking a fresh, single colony of Corvnebacterium glutamicum and growing a culture overnight in 10 mL CM (SM1) in a 250 mL shake flask at 30 degrees Celsius with agitation; (2) inoculating 200 mL of "Growth Media" with the overnight culture to an optical density (O.D.) of 660 nm of 0.1 in a 500 mL shake flask; (3) growing the culture at 30 degrees Celsius with agitation for 5-6 hours; (4) pouring the culture into a chilled, sealed, sterile 250 mL centrifuge bottle; Spin at 8-10K for ten minutes in Refrigerated Sorvall at 4 degrees Celsius; (5) pouring off the supernatant thoroughly and resuspending the cell pellet in an equal volume of ice-cold, sterile, deionized water; (6) centrifuging the sample again under the same conditions; (7) repeating the water wash remembering to keep everything ice-cold; (8) pouring off the supernatant thoroughly and resuspending the cell pellet in 1 mL of ice-cold, sterile 10% glycerol and transferring the cells to a chilled, sterile, 1.5 mL microcentrifuge tube; (9) spin the sample for 10 minutes in a refrigerated centrifuge; (10) pipetting off and discarding the supernatant, and resuspending the pellet in two to three times the pellet volume (200-400 µL) of 10% glycerol; and (11) aliquoting, if necessary, the cells into chilled tubes and freezing at -70 Celsius.

Plasmid DNAs are introduced into Corynebacterium glutamicum host cells by the following electroporation procedure: (1) pipetting 35 μ L, cell/glycerol solution onto the side wall of a chilled 0.1cm electrocuvette; (2) pipetting about 2-4 μ L of plasmid into the solution and mixing the sample by gentle pipetting up and down; (3) bringing the entire solution to the bottom of the electrocuvette by gentle tapping, avoiding the creation of bubbles; (4) keeping the sample on ice until ready for the electroshock step, wiping off any moisture on the outside of the electrocuvette prior to the electroshock administration, and shocking the cells one time at 1.5kV, 200Ω , 25μ F.

[0156] Cells are allowed to recover from electroporation by: (1) immediately pipetting 1 mL of warm "Recovery Media" into the electrocuvette and thoroughly mixing the solution by pipetting; (2) incubating the solution (in the electrocuvette) at 30 degrees Celsius for at least three hours for antibiotic resistance expression and cell recovery and (3) plating on selection media and incubating at 30 degrees Celsius for 3 days.

EXAMPLE 2

Preparation of L-Lysine Pathway Constructs Utilizing the Regulatory Regions of the Present Invention

[0157] Isolated nucleic acid molecules encoding L-lysine amino acid biosynthesis pathway genes are isolated by methods known to those of ordinary skill in the art.

[0158] Constructs which facilitate expression and regulation of L-lysine amino acid biosynthesis pathway genes are introduced into Corynebacterium strains by methods such as those described in Example 1. Any strain of Corynebacterium, particularly that of Corynebacterium glutamicum, may be utilized for the isolation of nucleic acid molecules for use in the preparation of vectors to improve amino acid biosynthetic pathway gene expression. Particularly preferred strains include: NRRL-B11474, ATCC 21799, ATCC 21529, ATCC 21543, and E12. As one

skilled in the art would know, the invention is not limited to these specific strain origins.

[0159] Methods and techniques common to the art of recombinant DNA technology are used in making vectors of the invention, as may be found in the many laboratory manuals cited and incorporated herein.

[0160] The polymerase chain reaction (PCR) technique may be used in the making of vectors of the invention. In a typical reaction, the standard 10X stock solution (100 mM Tris-HCL, pH 8.3, 500 mM KCL, 1.5 mM MgCl₂) is diluted to 1X for use. Typical reaction conditions were used for PCR amplication: 10 mM Tris, pH 8.3, 50 mM KCl, 1.5 mM MgCl₂, 0.01% gelatin, 200uM deoxynucleotides, 0.2-1.0uM primers and 2.0 U/100ul pfu polymerase. Standard cycling parameters were also employed in PCR reactions: For 30 cycles, template denaturation was performed at 94 °C for 30 sec.; 65 °C annealing temperature was performed for 1 min (or annealing temperature appropriate for particular primer pair); product extension was performed at 72 °C for 1 min (if product is <500 bp), 3 min (if product is >500 bp); and at the end of cycling, a final extension at 72 °C for 7 min was performed.

(a) Construction of pTrcLacZ

[0161] A plasmid construct designed to replicate as a plasmid in
Corynebacterium, which contains the β-galactosidase (lacZ) coding region operably associated with the lacl\(\frac{1}{2}\)/trc regulatory region, was constructed as follows. Plasmid pZap2, a plasmid which is capable of replicating in
Corynebacterium, was digested with Xbal and Smal, which linearize the plasmid just upstream of the lacZ gene. A cassette containing lacl\(\frac{1}{2}\) and the trc promoter
was excised from plasmid pTrc99A, available from Amersham Pharmacia
Biotech, by digestion with Sphl, filling in the overhang, and then digesting with
Xbal. The 1517-bp cassette (SEQ ID NO:30) was isolated from an agarose gel,
and ligated into the linearized pZap2 plasmid. The resulting ligation mix was

used to transform *Escherichia coli*, and colonies were isolated from LB plates supplemented with ampicillin ($100~\mu g/ml$), Xgal, and IPTG (LBXIA medium). Blue colonies were selected and plasmid isolates were verified by restriction digestion and electrophoresis.

(b) Construction of pAraBADLacZ

[0162] A plasmid construct designed to replicate as a plasmid in Corynebacterium, which contains the lacZ coding region operably associated with the araC/araBAD regulatory region, was constructed as follows. Plasmid pZap2 was digested with XbaI and SmaI as in (a). A cassette containing the araC gene and the pBAD promoter was excised from plasmid pBAD18-Cm, available from the American Type Culture Collection, Catalog No. 87396, by digestion with ClaI followed by a fill-in reaction and then digestion with XbaI. The cassette (i.e., nucleotides 1 to 1338 of SEQ ID NO:23) was isolated from an agarose gel and was ligated into the linearized pZap2 plasmid. Transformed E. coli colonies were plated on LBXIA medium as in (a), blue colonies were selected, and plasmid isolates were verified by restriction digestion and electrophoresis.

(c) Construction of pAceB10Zap2

[0163] A plasmid construct designed to replicate as a plasmid in
Corynebacterium, which contains the lacZ coding region operably associated
with the C. glutamicum aceB regulatory region, was constructed as follows. A
cassette containing the aceB promoter was PCR amplified from plasmid pSLD5
using forward primer 5' GTGCGGATCC TGGCTTTCCA ACGTTT 3' (SEQ ID
NO:31, BamHI site underlined) and reverse primer 5' CATGGG CTCGAGATGA
CCTGTGCCTA 3' (SEQ ID NO:32, SacI site underlined). The resulting PCR
product was digested with BamHI and SacI and ligated into the polylinker region
of cloning vector pD11 to produce pD11aceB. The resulting plasmid was

screened for orientation, and then the aceB promoter cassette was released from the vector by digestion with KpnI and XbaI and cloned into the KpnI and XbaI sites of plasmid pZap2 as described in (a). Transformed E. coli colonies were plated on LBXIA medium as in (a), blue colonies were selected, and plasmid isolates were verified by restriction digestion and electrophoresis.

(d) Construction of 131aceB/ask

chromosome, which contains the *C. glutamicum* aspartokinase (ask) coding region operably associated with the *C. glutamicum* aspartokinase (ask) coding region operably associated with the *C. glutamicum ace*B regulatory region, was constructed as follows. Plasmid pBGS131 (described in Spratt, B.G. et al. Gene 41:337-342 (1986)) was digested with *SmaI*. The aceB cassette was released from plasmid pD11aceB, produced as in (c), by digestion with appropriate restriction enzymes and the cassette was subjected to a fill-in reaction to produce blunt ends. The cassette was then ligated into the digested pBGS131. Plasmid isolates were tested for the proper orientation of the aceB promoter by restriction digestion and electrophoresis, resulting in plasmid 131aceB. The *C. glutamicum ask* coding region was released from pTrc99Aask (produced as in (f), infra) by digestion with SacI and KpnI and cloned into the SacI and KpnI sites of 131aceB, to produce 131aceBask.

(e) Construction of PD10aceBask

[0165] A plasmid construct designed to replicate as a plasmid in C. glutamicum, which contains the C. glutamicum aspartokinase (ask) coding region operably associated with the C. glutamicum aceB regulatory region, was constructed as follows. A cassette containing the ask coding region operably associated with the aceB regulatory region was released from 131aceBask, produced as in (d), by digestion with BamHI and KpnI. The resulting fragment was cloned into the

BamHI and KpnI sites of plasmid pD10, which is capable of replicating in C. glutamicum, to produce pD10aceBask.

(f) Construction of 131LacI^q/trc-ask

[0166] A plasmid construct designed to integrate into the *C. glutamicum* chromosome, which contains the *C. glutamicum* aspartokinase (ask) coding region operably associated with the lacl⁹/trc regulatory region, was constructed as follows. The PCR-amplified ask coding region was digested with SacI and KpnI, and ligated into plasmid pTrc99A which had been digested with SacI and KpnI to produce pTrc99Aask. A cassette containing the ask coding region operably associated with the lacl⁹/trc regulatory region was then released from pTrc99Aask by digestion with SphI, and was cloned into the SphI site of plasmid pBGS131 to produce 131Lacl⁹/trc-ask.

(g) Construction of Additional LacZ Plasmid Expression Constructs

[0167] Plasmid constructs designed to replicate as a plasmid in Corynebacterium, which contain the lacZ coding region operably associated with C. glutamicum regulatory regions contained in SEQ ID NOs 1, 2, and 4-22, are constructed as follows. Cassettes containing the regulatory regions present in SEQ ID NOs 1, 2, and 4-22 are PCR amplified from the C. glutamicum chromosome using primers engineered to introduce appropriate restriction enzyme recognition sites onto the ends of the cassettes. In designing the primers, care is taken to introduce sites for restriction enzymes which will not digest the cassette internally. The resulting PCR products are digested with the chosen restriction enzymes, and the fragments are ligated into a polylinker region of a standard cloning vector. The resulting plasmids are screened for orientation, and then the cassettes are released from the vector by digestion with appropriate restriction enzymes, filled in as necessary, and cloned into the polylinker region upstream of the lacZ gene in

plasmid pZap2 as described in (a). Transformed *E. coli* colonies are plated on LBXIA medium as in (a), blue colonies are selected, and plasmid isolates are verified by restriction digestion and electrophoresis.

(h) Construction Additional Aspartokinase Expression Constructs Designed to Integrate

Plasmid constructs designed to integrate into the C. glutamicum [0168] chromosome, which contain the C. glutamicum aspartokinase (ask) coding region operably associated with C. glutamicum regulatory regions contained in SEQ ID NOs 1, 2, and 4-22, are constructed as follows. Plasmid pBGS131 is digested with Smal, or other appropriate restriction enzyme. The regulatory cassettes prepared as in (g) are released by digestion with appropriate restriction enzymes and the cassettes are subjected to fill-in reactions to produce blunt ends as needed. The cassettes are then ligated into the digested pBGS131. Plasmid isolates are tested for the proper orientation of the regulatory regions by restriction digestion and electrophoresis, to produce plasmids 131SEQ ID NO:1, 131 SEQ ID NO:2, and 131SEQ ID NO:4-131 SEQ ID NO:22. The C. glutamicum ask coding region is released from pTrc99Aask (produced as in (f)) by digestion with SacI and KpnI (or other appropriate restriction enzymes), or is PCR amplified from a C. glutamicum chromosome, and is cloned into the corresponding sites of plasmids 131SEQ ID NO:1, 131 SEQ ID NO:2, and 131SEQ ID NO:4-131 SEQ ID NO:22, to produce the plasmids 131SEQ ID NO:1-ask, 131 SEQ ID NO:2-ask, and 131SEQ ID NO:4-ask through 131 SEQ ID NO:22-ask.

- (i) Construction of Additional Aspartokinase Constructs Designed to Replicate as a Plasmid
- [0169] Plasmid constructs designed to replicate as a plasmid in C. glutamicum, which contain the C. glutamicum aspartokinase (ask) coding region operably associated with C. glutamicum regulatory regions contained in SEQ ID NO1,2,

and 4-22, are constructed as follows. Cassettes containing the ask coding region operably associated with the regulatory regions contained in SEQ ID NO:1, SEQ ID NO:2, and SEQ ID NO:4-SEQ ID NO:22 are released from plasmids 131 SEQ ID NO:1-ask, 131 SEQ ID NO:2-ask, and 131 SEQ ID NO:4-ask through 131 SEQ ID NO:22-ask, produced as in (h), by digestion with appropriate restriction enzymes. The resulting fragments are cloned into corresponding sites of plasmid pD10, which is capable of replicating in C. glutamicum, to produce plasmids PD10/SEQ ID NO:1-ask, PD10/SEQ ID NO:2-ask, and PD10/SEQ ID NO:4-ask through PD10/SEQ ID NO:2-ask.

EXAMPLE 3

Use of LacZ as a Reporter to Demonstrate Transcriptional Regulation by araBAD (SEQ ID NO:23) and the lacI⁹/trc Transcriptional Complex (SEQ ID NO:30) in Corvnebacterium (41-004002)

[0170] The ability of the heterologous transcriptional regulatory regions lac[¹/trc and araBAD (with the araC repressor) to control gene expression in C. glutamicum was tested by measuring expression of β-galactosidase. Plasmids pTrcLacZ and pAraBADLacZ, produced as in Examples 2(a) and 2(b), respectively, were transferred into C. glutamicum according to the methods described in Example 1. The lacZ reporter gene, regulated by either of the two regulatory regions, was expressed from a plasmid replicating in C. glutamicum, grown in the presence of an inducer, IPTG in the case of the lacI¹/trc regulatory region, or arabinose in the case of the araC/araBAD regulatory region. The level of β-galactosidase production was measured, and the results are shown in Table 3.

TABLE 3

regulator/promoter	Inducer	nmol/min/mg	
		Mean	S.D.
none	none	0.40	0.14
LacI ^q /trc	none	3.84	0.57
LacI ^q /trc	+IPTG (30mg/L)	77.23	0.32
AraC/AraBAD	none	0.47	0.49
AraC/AraBAD	+arabinose (1 g/L)	5.51	0.46

[0171] The trc regulated construct shows uninduced activity. The araBAD regulated construct showed no uninduced activity and responded to arabinose. The araBAD promoter resulted in a β-galactosidase expression level that was 14-fold lower than that observed with the trc promoter in this organism.

EXAMPLE 4

Use of β -Galactosidase Activity to Demonstrate the Response of the C. glutamicum aceB Promoter to Acetate, Glucose and Ethanol (98-0040002)

[0172] The ability of the C. glutamicum aceB regulatory region (SEQ ID NO:3) to control gene expression of a heterologous reporter gene in C. glutamicum was tested by measuring expression of β-galactosidase from plasmid pAceB10Zap2, produced as in Example 2(c). In this construct, the lacZ reporter gene, regulated by aceB, is expressed from a plasmid replicating in C. glutamicum. Response of the regulatory region to various inducers was tested. The results are shown in Table 4

TABLE 4

Rich Medium	nmol/min/mg
Minus sucrose	10
+ 2% glucose	3
+ 0.2% ammonium acetate	26
+ 2% ammonium acetate	50
+ 2% glucose + 0.2% ammonium acetate	7
+ 2% glucose + 2% ammonium acetate	15
+ 2% ethanol	28

[0173] Glucose repressed reporter activity. Ethanol induced activity but not as strongly as acetate. The uninduced control also showed activity implying that aceB is self-regulated in this medium.

EXAMPLE 5

Use of the trc Promoter to Regulate Aspartokinase Activity when Integrated into the Chromosome of C. glutamicum (90-004002)

[0174] The ability of the lacl^q/trc (SEQ ID NO:30) regulatory region to control expression of aspartokinase, an enzyme in the *C. glutamicum* L-lysine biosynthesis pathway (encoded by the *ask* gene), was tested as follows. The 131-2 and 131-5 *C. glutamicum* strains have the lacl^q/trc transcriptional regulatory region, in operable association with the *ask* gene, integrated into the *C. glutamicum* chromosome. These strains were prepared by introducing plasmid 131Lacl^q/trc-ask into *C. glutamicum* by the methods described in Example 1. The level of aspartokinase activity was measured both with and without the addition of the inducer IPTG. The results are shown in Table 5.

TABLE 5

Strain	Regulator/promoter-	Inducer	nmol/min/mg
	gene		
131-2	lacIq/trc-ask	none	59
131-2	lacI4/trc-ask	+IPTG (30 mg/L)	117
131-5	lacF/trc-ask	none	59
131-5	lacIq/trc-ask	+IPTG (30 mg/L)	123

[0175] The level of aspartokinase activity is doubled upon induction when the trc regulon is integrated into the chromosome.

EXAMPLE 6

Lysine Production by Integrated, Uninduced lac I⁴/trc-ask Constructs (84-004002)

[0176] This example shows the influence on lysine production of the uninduced lacl*/trc-ask construct when integrated into the chromosome of C. glutamicum. The lacf*/trc regulatory region was used to control expression of aspartokinase (encoded by the ask gene), an enzyme in the C. glutamicum L-lysine biosynthesis pathway. Bacterial cultures of 131-2 and 131-5, prepared as described in Example 5, were grown in shake flasks. BF100 is C. glutamicum strain which is a high level lysine producer. The level of aspartokinase activity was measured without the addition of an inducer. The results are shown in Table 6.

TABLE 6

Strain	Induction	O.D.	Titre	Yield	S.P.	
BF100	none	43	23	39	53	
131-2	none	34	27	46	82	
131-5	none	35	28	47	82	
O.D. = op	tical density a	t 660nm				
Titre = grams Lysine/liter						
Yield = grams lysine made/grams dextrose consumed						
S.P. = grams lysine/O.D.						

[0177] The leakiness of the trc promoter in this organism may be responsible for the improved productivity in the absence of an inducer.

EXAMPLE 7

Improvement of Lysine Production in Shake Flasks by aceB-Aspartokinase Constructs with No Exogenous Induction (84-0040002)

Aspartokinase construct when integrated into the *C. glutamicum* chromosome.

The *C. glutamicum ace*B regulatory region (SEQ ID NO:3) was used to control expression of aspartokinase (encoded by the *ask* gene), an enzyme in the *C. glutamicum* L-lysine biosynthesis pathway. Bacterial cultures were grown in shake flasks. Strain 131-6 is a *C. glutamicum* strain which has the *ace*B transcriptional regulatory region in operable association with the *ask* gene, integrated into the bacterial chromosome. Strain 131-6 was prepared by introducing plasmid 131*ace*B/*ask*, produced as described in Example 2(d), into *C. glutamicum* by the methods described in Example 1. BF100 is *C. glutamicum* strain which is a high level lysine producer. The level of aspartokinase activity was measured either with or without the addition of the inducer acetate. The results are shown in Table 7.

TABLE 7

Strain	Induction	O.D.	Titre	Yield
BF100	none	43	25	37
	acetate	42	24	36
131-6	none	50	27	41
	acetate	47	27	40

Titer = g lysine/L medium

Yield = g lysine/ g glucose consumed * 100%

[0179] Induction by ammonium acetate is not required to observe an improvement in lysine production by this construct.

EXAMPLE 8

Improvement of Lysine Production in Fermentors by aceB-aspartokinase Constructs with No Exogenous Induction (Lys002 Lys006RLK)

[0180] This example shows the influence on lysine production of uninduced aceB-Aspartokinase constructs which are expressed from a plasmid (PD10aceBask) replicating in C. glutamicum, or which are integrated

[0181] into the *C. glutamicum* chromosome as in strain 131-6, described in Example 7. The *C. glutamicum aceB* regulatory region (SEQ ID NO:3) was used to control expression of aspartokinase (encoded by the *ask* gene), an enzyme in the *C. glutamicum* L-lysine biosynthesis pathway. Bacterial cultures were grown in fermentors. The *C. glutamicum* 131-*aceB*-aspartokinase strain, 131-6, has the *aceB* transcriptional regulatory region in operable association with the *ask* gene, integrated into the *C. glutamicum* chromosome. The *C. glutamicum* PD10-*aceB*-aspartokinase strain has the *aceB* transcriptional regulatory region in operable association with the *ask* gene, expressed from a plasmid (PD10*aceBask*, produced as described in Example 2(e)) replicating in *C. glutamicum*. BF100 is *C. glutamicum* strain which is a high level lysine producer. Aspartokinase activity was measured in each of these three strains. The results are shown in Table 8.

TABLE 8

Strain	Titer	Yield	Product
BF100	127	32.9	735
131-aceB-aspartokinase	129	39.8	833
BF100	138	40.7	769
PD10-aceB-aspartokinase	135	45.8	830

Titer = g lysine/L medium

Yield = g lysine/g glucose consumed * 100%

Product = g lysine made in that fermented

[0182] The aceB-aspartokinase constructs contributed to productivity whether they were maintained on a plasmid or integrated into the chromosome. In neither case did the constructs need to be induced by an exogenous compound.

EXAMPLE 9

Influence of Medium on Beta Galactosidase Activity Using Various Use of LacZ as a Reporter to Demonstrate Transcriptional Regulation by the Regulatory Regions Contained in SEQ ID NOs 1,2, and 4-22 in Corynebacterium

[0183] The ability of certain transcriptional regulatory regions disclosed herein to control gene expression in *C. glutamicum* was tested by measuring expression of β-galactosidase. The promoters used include (SEQ ID NO: 27) Tac, (SEQ ID NO: 1) Pta, (SEQ ID NO: 2) AceA, (SEQ ID NO: 3) AceB, (SEQ ID NO: 4) Adh, (SEQ ID NO: 5) AldB, (SEQ ID NO: 6) PoxB, (SEQ ID NO: 7) Ldh, (SEQ ID NO: 9) MalZ, (SEQ ID NO: 10) BglX, (SEQ ID NO: 13) HisD, (SEQ ID NO: 14) PyrR, (SEQ ID NO: 15) PurD, (SEQ ID NO: 18) DnaK, (SEQ ID NO: 20) GrpE, (SEQ ID NO: 21) ClpB, (SEQ ID NO: 33) LeuA, (SEQ ID NO: 34) IlvA and (SEQ ID NO: 35) IlvB. The reporter gene was expressed from plasmids replicating in *C. glutamicum*, and was driven by the regulatory regions contained in SEQ ID NOs 1,2, and 4-22. The plasmids were prepared as described in

Example 2(g). Increased expression of beta-galactosidase under the transcriptional control of these transcriptional regulatory regions is shown in Table 9. The "Seed" Medium is experimentally determined to support high growth. The "Main" Medium is formulated to induce higher expression of products, such as lysine, for example. Exemplary medias were described *supra*.

[0184] For certain regulatory regions, β-galactosidase levels are measured without the addition or any inducer or in the presence of various inducers, e.g., acetic acid, pyruvate, ethanol, a starch subunit, a sugar, e.g., fructose, maltose, lactose or arabinose, a cellulose subunit, a fatty acid, or a triglyceride. In addition, certain regulatory regions are tested by growth with a suppressor, e.g., a purine, a pyrimidine, and amino acid, or oxygen, followed by a shift to medium lacking the suppressor at which time β-galactosidase levels were measured. For certain of the transcriptional regulatory regions, β-galactosidase levels are measured at optimal growth temperatures, and at temperatures either greater than or less than the optimal growth temperature. The β-galactosidase levels were measured using certain transcriptional regulatory regions at varying temperatures as shown in Table 10.

TABLE 9

Influence of Medium on beta-galactosidase Reporter Activity

(units = nmol/min/mg protein)

		<u>Medium</u>		
SEQ ID NO	Promoter	Seed(16hr)	Main(48hr)	Change
	Promoterless	2	20	19
27	Tac	231	342	111
1	Pta	134	111	-23
2	AceA	0	63	63
3	AceB	0	25	25
4	Adh	0	65	65

5	AldB	9	260	251
6	PoxB	150	276	126
7	Ldh	205	291	86
9	MalZ	62	234	172
10	BglX	57	208	151
13	HisD	119	241	122
14	PyrR	161	289	128
15	PurD	9	56	47
18	DnaK	61	223	162
20	GrpE	40	149	109
21	ClpB	104	233	129
33	LeuA	174	267	193
34	IlvA	120	229	109
35	llvB	22	166	144

TABLE 10

Effect of Temperature Change on beta-galactosidase Reporter Activity

(units = nmol/min/mg protein; mean (standard deviation))

temperature shift in seed medium for 4 hrs

SEQ ID NO	Promoter	30°C	36°C	40°C
	Promoterless	0(0.7)	0(0.2)	0(0.7)
21	clpB	76(7.3)	123(2.5)	165(1.6)
18	dnaK	70(0.8)	119(3.1)	169(4.5)
20	grpE	47(1.6)	107(1.6)	97(2.6)

All 3 promoters increased reporter activity at 36°C

The grpE promoter doubled the activity at both 36 °C and 40 °C

All 3 promoters increased reporter activity at 40°C

[0185]

EXAMPLE 10

Improvement of Lysine Production Through Transcriptional Regulation of Aspartokinase by Regulatory Regions Contained in SEQ ID NOs 1,2, and 4-22

This example shows the influence on lysine production of induced or uninduced constructs in which the transcriptional regulatory regions of SEQ ID NOs 1,2, and 4-22 are used to express aspartokinase, either from a plasmid replicating in C. glutamicum, or when integrated into the C. glutamicum chromosome. The various C. glutamicum regulatory regions are used to control expression of aspartokinase, an enzyme in the C. glutamicum L-lysine biosynthesis pathway. Bacterial cultures are grown either in shake flasks or fermentors. C. glutamicum strains are prepared in which the transcriptional regulatory regions of SEQ ID NOs 1,2, and 4-22, in operable association with the ask gene, are integrated into the C. glutamicum chromosome. These strains are prepared by introducing plasmids 131 SEQ ID NO:1-ask, 131 SEQ ID NO:2-ask, and 131SEQ ID NO:4-ask through 131 SEQ ID NO:22-ask, prepared as described in Example 2(h), into C. glutamicum by methods described in Example 1. Additional strains, with self-replicating plasmids are prepared by introducing plasmids PD10/SEQ ID NO:1-ask, PD10/SEQ ID NO:2-ask, and PD10/SEQ ID NO:4-ask through PD10/SEQ ID NO:22-ask, produced as described in Example 2(i), into C. glutamicum by methods described in Example 1. BF100, a high level lysine producer, is used as a control. Aspartokinase activity is measured in each of the resulting strains. For certain regulatory regions, aspartokinase levels are measured without the addition or any inducer, or in the presence of various inducers, e.g., acetic acid, pyruvate, ethanol, a starch subunit, a sugar, e.g., fructose, maltose, lactose or arabinose, a cellulose subunit, a fatty acid, or a triglyceride. In addition, certain regulatory regions are tested by growth either without, or with a suppressor, e.g., a purine, a pyrimidine, and amino acid, or oxygen, followed by a shift to medium lacking the suppressor at which time aspartokinase levels are measured. Finally, for certain regulatory regions, aspartokinase levels are measured at optimal growth temperatures, and also at temperatures either greater than or less than the optimal growth temperature.

* * * * *

[0186] Having now fully described the present invention in some detail by way of illustration and example for purposes of clarity of understanding, it will be obvious to one of ordinary skill in the art that same can be performed by modifying or changing the invention with a wide and equivalent range of conditions, formulations and other parameters thereof, and that such modifications or changes are intended to be encompassed within the scope of the appended claims.

[0187] All publications, patents and patent applications mentioned in this specification are indicative of the level of skill of those skilled in the art to which this invention pertains, and are herein incorporated by reference to the same extent as if each individual publication, patent or patent application was specifically and individually indicated to be incorporated by reference.